# (19) World Intellectual Property Organization International Bureau





# (43) International Publication Date 5 April 2001 (05.04.2001)

#### **PCT**

# (10) International Publication Number WO 01/22920 A2

(51) International Patent Classification7:

A61K

- (21) International Application Number: PCT/US00/26524
- (22) International Filing Date:

28 September 2000 (28.09.2000)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

60/157,137 29 September 1999 (29.09.1999) US 60/163,280 3 November 1999 (03.11.1999) US

- (71) Applicant (for all designated States except US): HUMAN GENOME SCIENCES, INC. [US/US]; 9410 Key West Avenue, Rockville, MD 20850 (US).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): RUBEN, Steven, M. [US/US]; 18528 Heritage Hills Drive, Olney, MD 20832 (US). BARASH, Steven, C. [US/US]; 111 Watkins Pond Blvd., #301, Rockville, MD 20850 (US). BIRSE, Charles, E. [GB/US]; 13822 Saddleview Drive, North Potomac, MD 20878 (US). ROSEN, Craig, A. [US/US]; 22400 Rolling Hill Road, Laytonsville, MD 20882 (US).

- (74) Agents: HOOVER, Kenley, K. et al.; c/o Human Genome Sciences, Inc., 9410 Key West Avenue, Rockville, MD 20850 (US).
- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

#### Published:

 Without international search report and to be republished upon receipt of that report.

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.



A2

#### (54) Title: COLON AND COLON CANCER ASSOCIATED POLYNUCLEOTIDES AND POLYPEPTIDES

(57) Abstract: This invention relates to newly identified colon or colon cancer related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "colon cancer antigens", and the use of such colon cancer antigens for targeting specific cell types and/or diagnosing, detecting, preventing and treating disorders of the colon, particularly the presence of colon cancer and colon cancer metastases. This invention relates to colon cancer antigens as well as vectors, host cells, antibodies directed to colon cancer antigens and the recombinant or synthetic methods for producing the same. Also provided are diagnostic methods for diagnosing and treating, preventing and/or prognosing disorders related to the colon, including colon cancer, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of colon cancer antigens of the invention. The present invention further relates to inhibiting the production and function of the polypeptides of the present invention.

WO 01/22920 PCT/US00/26524

# Colon and Colon Cancer Associated Polynucleotides and Polypeptides

#### Field of the Invention

5

10

15

This invention relates to newly identified colon or colon cancer related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "colon cancer antigens," and the use of such colon cancer antigens for targeting specific cell types and/or diagnosing, detecting, preventing and treating disorders disorders of the colon, particularly the presence of colon cancer and colon cancer metastases. This invention relates to colon cancer antigens as well as vectors, host cells, antibodies directed to colon cancer antigens and the recombinant or synthetic methods for producing the same. Also provided are diagnostic methods for diagnosing and treating, preventing and/or prognosing disorders related to the colon, including colon cancer, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of colon cancer antigens of the invention. The present invention further relates to inhibiting the production and function of the polypeptides of the present invention.

## Background of the Invention

20

Cell growth is a carefully regulated process which responds to specific needs of the body. Occasionally, the intricate, and highly regulated controls dictating the rules for cellular division break down. When this occurs, the cell begins to grow and divide independently of its homeostatic regulation resulting in a condition commonly referred to as cancer. In fact, cancer is the second leading cause of death among Americans aged 25-44.

25

30

Colorectal cancers are among the most common cancers in men and women in the U.S. and are one of the leading causes of death. Other than surgical resection no other systemic or adjuvant therapy is available. Vogelstein and colleagues have described the sequence of genetic events that appear to be associated with the multistep process of colon cancer development in humans (Trends Genet 9(4):138-41 (1993)). An understanding of the molecular genetics of carcinogenesis, however, has not led to preventative or therapeutic measures. It can be expected that advances in molecular genetics will lead to better risk

WO 01/22920 PCT/US00/26524

2

assessment and early diagnosis but colorectal cancers will remain a deadly disease for a majority of patients due to the lack of an adjuvant therapy. Adjuvant or systemic treatments are likely to arise from a better understanding of the autocrine factors responsible for the continued proliferation of cancer cells.

Colorectal carcinoma is a malignant neoplastic disease. There is a high incidence of colorectal carcinoma in the Western world, particularly in the United States. Tumors of this type often metastasize through lymphatic and vascular channels. Many patients with colorectal carcinoma eventually die from this disease. In fact, it is estimated that 62,000 persons in the United States alone die of colorectal carcinoma annually.

At the present time the only systemic treatment available for colon cancer is chemotherapy. However, chemotherapy has not proven to be very effective for the treatment of colon cancers for several reasons, the most important of which is the fact that colon cancers express high levels of the MDR gene (that codes for multi-drug resistance gene products). The MDR gene products actively transport the toxic substances out of the cell before the chemotherapeutic agents can damage the DNA machinery of the cell. These toxic substances harm the normal cell populations more than they harm the colon cancer cells for the above reasons.

There is no effective systemic treatment for treating colon cancers other than surgically removing the cancers. In the case of several other cancers, including breast cancers, the knowledge of growth promoting factors (such as EGF, estradiol, IGF-11) that appear to be expressed or effect the growth of the cancer cells, has been translated for treatment purposes. But in the case of colon cancers this knowledge has not been applied and therefore the treatment outcome for colon cancers remains bleak.

Thus, the discovery of new human colon and colon cancer related polynucleotides and the polypeptides encoded by them satisfies a need in the art by providing new compositions which are useful in the diagnosis, prevention and treatment of disorders of the colon, particularly tumors, especially of the intestine; inflammatory disorders; enterocolitis; miscellaneous intestinal inflammatory disorders; ulcerative disorders; and/or noncancerous tumors.

5

10

15

20

25

3

## Summary of the Invention

This invention relates to newly identified colon and colon cancer related polynucleotides and the polypeptides encoded by these polynucleotides herein collectively known as "colon cancer antigens." This invention relates to colon and colon cancer related polypeptides as well as vectors, host cells, antibodies directed to colon cancer antigens and the recombinant methods for producing the same. Also provided are diagnostic methods for diagnosing and treating, preventing and/or prognosing disorders related to the colon, including colon cancer, and therapeutic methods for treating such disorders. The invention further relates to screening methods for identifying agonists and antagonists of colon cancer antigens of the invention.

## **Detailed Description**

#### **Tables**

5

10

15

20

25

30

Table 1 summarizes some of the colon cancer antigens encompassed by the invention (including cDNA clones related to the sequences (Clone ID NO:Z), polynucleotide sequences (contig identifier (Contig ID:) or sequence identifier (Sequence ID:) and nucleotide sequence identifier (SEQ ID NO:X)) and further summarizes certain characteristics of the colon and colon cancer related polynucleotides and the polypeptides encoded thereby. The first column shows the "SEQ ID NO:X" for each of the 4277 colon and colon cancer related polynucleotide sequences of the invention. The second column provides a unique "Sequence/Contig ID" identification for each of the colon and colon cancer related polynucleotide and/or polypeptide sequences. The third column, "Gene Name," and the fourth column, "Overlap," provide a putative identification of the gene based on the sequence similarity of its translation product to an amino acid sequence found in a publicly accessible gene database and the database accession no. for the database sequence having similarity, respectively. The sixth and seventh columns provide the location (nucleotide position nos. within the sequence/contig), "Start" and "End", in the polynucleotide sequence "SEQ ID NO:X" that delineate the preferred ORF shown in the sequence listing as SEQ ID NO:Y (column five). The eighth and ninth columns provide the "%Id" (percent identity) and "%Si" (percent similarity), respectively, observed between the aligned sequence segments of the translation product of SEQ ID NO:X and the database sequence. The tenth column provides a

5

10

15

20

25

30

4

unique "Clone ID:Z" for a cDNA clone related to each contig sequence. The eleventh column provides the "Cloning vector" contained in the cDNA clone ID.

PCT/US00/26524

Table 2 summarizes ATCC Deposits, Deposit dates, and ATCC designation numbers of deposits made with the ATCC in connection with the present application.

Table 3 summarizes the expression profile of polynucleotides corresponding to the clones disclosed in Table 1. The first column provides a unique clone identifier, "Clone ID", for a cDNA clone related to each contig sequence disclosed in Table 1. Column 2, "Library Codes" shows the expression profile of tissue and/or cell line libraries which express the polynucleotides of the invention. Each Library Code in column 2 represents a tissue/cell source identifier code corresponding to the Library Code and Library description provided in Table 5. Expression of these polynucleotides was not observed in the other tissues and/or cell libraries tested. One of skill in the art could routinely use this information to identify tissues which show a predominant expression pattern of the corresponding polynucleotide of the invention or to identify polynucleotides which show predominant and/or specific tissue expression.

Table 4, column 1, provides a nucleotide sequence identifier, "SEQ ID NO:X," that matches a nucleotide SEQ ID NO:X disclosed in Table 1, column 5. Table 4, column 2, provides the chromosomal location, "Cytologic Band or Chromosome," of polynucleotides corresponding to SEQ ID NO:X. Chromosomal location was determined by finding exact matches to EST and cDNA sequences contained in the NCBI (National Center for Biotechnology Information) UniGene database. Given a presumptive chromosomal location, disease locus association was determined by comparison with the Morbid Map, derived from Online Mendelian Inheritance in Man (Online Mendelian Inheritance in Man, OMIMTM. McKusick-Nathans Institute for Genetic Medicine, Johns Hopkins University (Baltimore, MD) and National Center for Biotechnology Information, National Library of Medicine (Bethesda, MD) 2000. World Wide Web URL: http://www.ncbi.nlm.nih.gov/omim/). If the putative chromosomal location of the Query overlapped with the chromosomal location of a Morbid Map entry, the OMIM reference identification number of the morbid map entry is provided in Table 4, column 3, labelled "OMIM ID." A key to the OMIM reference identification numbers is provided in Table 6.

WO 01/22920 PCT/US00/26524

Table 5 provides a key to the Library Code disclosed in Table 3. Column 1 provides the Library Code disclosed in Table 3, column 2. Column 2 provides a description of the tissue or cell source from which the corresponding library was derived.

5

Table 6 provides a key to the OMIM reference identification numbers disclosed in Table 4, column 3. OMIM reference identification numbers (Column 1) were derived from Online Mendelian Inheritance in Man (Online Mendelian Inheritance in Man, OMIM. McKusick-Nathans Institute for Genetic Medicine, Johns Hopkins University (Baltimore, MD) and National Center for Biotechnology Information, National Library of Medicine, (Bethesda, MD) 2000. World Wide Web URL: http://www.ncbi.nlm.nih.gov/omim/). Column 2 provides diseases associated with the cytologic band disclosed in Table 4, column 2, as determined using the Morbid Map database.

Table 7 indicates public ESTs, of which at least one, two, three, four, five, ten, fifteen or more of any one or more of these public EST sequences are optionally excluded from certain embodiments of the invention.

Table 8 lists residues comprising antigenic epitopes of antigenic epitope-bearing fragments present in each of the preferred ORFs (SEQ ID NO:Y) encoded by the colon or colon cancer related polynucleotides described in Table 1 as predicted by the inventors using the algorithm of Jameson and Wolf, (1988) Comp. Appl. Biosci. 4:181-186. The Jameson-Wolf antigenic analysis was performed using the computer program PROTEAN (Version 3.11 for the Power MacIntosh, DNASTAR, Inc., 1228 South Park Street Madison, WI). Colon and/or colon cancer related polypeptides shown in Table 1 may possess one or more antigenic epitopes comprising residues described in Table 8. It will be appreciated that depending on the analytical criteria used to predict antigenic determinants, the exact address of the determinant may vary slightly. The residues and locations shown described in Table 8 correspond to the amino acid sequences for each colon and/or colon cancer related polypeptide sequence shown in the Sequence Listing.

Table 9 shows the cDNA libraries sequenced, and ATCC designation numbers and vector information relating to these cDNA libraries.

#### **Definitions**

5

10

15

20

25

30

The following definitions are provided to facilitate understanding of certain terms used throughout this specification.

5

10

15

20

25

30

6

PCT/US00/26524

In the present invention, "isolated" refers to material removed from its original environment (e.g., the natural environment if it is naturally occurring), and thus is altered "by the hand of man" from its natural state. For example, an isolated polynucleotide could be part of a vector or a composition of matter, or could be contained within a cell, and still be "isolated" because that vector, composition of matter, or particular cell is not the original environment of the polynucleotide. The term "isolated" does not refer to genomic or cDNA libraries, whole cell total or mRNA preparations, genomic DNA preparations (including those separated by electrophoresis and transferred onto blots), sheared whole cell genomic DNA preparations or other compositions where the art demonstrates no distinguishing features of the polynucleotide/sequences of the present invention.

As used herein, a "polynucleotide" refers to a molecule having a nucleic acid sequence contained in SEQ ID NO:X or the cDNA clone deposited with the ATCC. For example, the polynucleotide can contain the nucleotide sequence of the full length cDNA sequence, including the 5' and 3' untranslated sequences, the coding region, as well as fragments, epitopes, domains, and variants of the nucleic acid sequence. Moreover, as used herein, a "polypeptide" refers to a molecule having the translated amino acid sequence generated from the polynucleotide as broadly defined.

In the present invention, "SEQ ID NO:X" was often generated by overlapping sequences contained in multiple clones (contig analysis). A representative clone containing all or most of the sequence for SEQ ID NO:X is deposited at Human Genome Sciences, Inc. (HGS) in a catalogued and archived library. As shown in Table 1, each clone is identified by a cDNA Clone ID (Identifier). Each Clone ID is unique to an individual clone and the Clone ID is all the information needed to retrieve a given clone from the HGS library. In addition to the individual cDNA clone deposits, the cDNA clones were deposited at the American Type Culture Collection (hereinafter "ATCC"). As mentioned below, Table 1 correlates the Clone ID names with SEQ ID NOs. Furthermore, it is possible to retrieve a given cDNA clone from the ATCC deposit by techniques known in the art and described elsewhere herein. The ATCC is located at 10801 University Boulevard, Manassas, Virginia 20110-2209, USA. The ATCC deposits were made persuant to the terms of the Budapest Treaty on the international recognition of the deposit of microorganisms for the purposes of patent procedure.

5

10

15

20

25

30

7

PCT/US00/26524

In specific embodiments, the polynucleotides of the invention are at least 15, at least 30, at least 50, at least 100, at least 125, at least 500, or at least 1000 continuous nucleotides but are less than or equal to 300 kb, 200 kb, 100 kb, 50 kb, 15 kb, 10 kb, 7.5kb, 5 kb, 2.5 kb, 2.0 kb, or 1 kb, in length. In a further embodiment, polynucleotides of the invention comprise a portion of the coding sequences, as disclosed herein, but do not comprise all or a portion of any intron. In another embodiment, the polynucleotides comprising coding sequences do not contain coding sequences of a genomic flanking gene (i.e., 5' or 3' to the gene of interest in the genome). In other embodiments, the polynucleotides of the invention do not contain the coding sequence of more than 1000, 500, 250, 100, 50, 25, 20, 15, 10, 5, 4, 3, 2, or 1 genomic flanking gene(s).

A "polynucleotide" of the present invention also includes those polynucleotides capable of hybridizing, under stringent hybridization conditions, to sequences contained in SEQ ID NO:X, the complement thereof, or the cDNA clone within the pool of cDNA clones deposited with the ATCC. "Stringent hybridization conditions" refers to an overnight incubation at 42 degree C in a solution comprising 50% formamide, 5x SSC (750 mM NaCl, 75 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5x Denhardt's solution, 10% dextran sulfate, and 20 μg/ml denatured, sheared salmon sperm DNA, followed by washing the filters in 0.1x SSC at about 65 degree C.

Also contemplated are nucleic acid molecules that hybridize to the polynucleotides of the present invention at lower stringency hybridization conditions. Changes in the stringency of hybridization and signal detection are primarily accomplished through the manipulation of formamide concentration (lower percentages of formamide result in lowered stringency); salt conditions, or temperature. For example, lower stringency conditions include an overnight incubation at 37 degree C in a solution comprising 6X SSPE (20X SSPE = 3M NaCl; 0.2M NaH<sub>2</sub>PO<sub>4</sub>; 0.02M EDTA, pH 7.4), 0.5% SDS, 30% formamide, 100 ug/ml salmon sperm blocking DNA; followed by washes at 50 degree C with 1XSSPE, 0.1% SDS. In addition, to achieve even lower stringency, washes performed following stringent hybridization can be done at higher salt concentrations (e.g. 5X SSC).

Note that variations in the above conditions may be accomplished through the inclusion and/or substitution of alternate blocking reagents used to suppress background in hybridization experiments. Typical blocking reagents include Denhardt's reagent, BLOTTO, heparin, denatured salmon sperm DNA, and commercially available proprietary formulations.

5

10

15

20

25

30

8

The inclusion of specific blocking reagents may require modification of the hybridization conditions described above, due to problems with compatibility.

PCT/US00/26524

Of course, a polynucleotide which hybridizes only to polyA+ sequences (such as any 3' terminal polyA+ tract of a cDNA shown in the sequence listing), or to a complementary stretch of T (or U) residues, would not be included in the definition of "polynucleotide," since such a polynucleotide would hybridize to any nucleic acid molecule containing a poly (A) stretch or the complement thereof (e.g., practically any double-stranded cDNA clone generated using oligo dT as a primer).

The polynucleotide of the present invention can be composed of any polyribonucleotide or polydeoxribonucleotide, which may be unmodified RNA or DNA or modified RNA or DNA. For example, polynucleotides can be composed of single- and double-stranded DNA, DNA that is a mixture of single- and double-stranded regions, single- and double-stranded RNA, and RNA that is mixture of single- and double-stranded regions, hybrid molecules comprising DNA and RNA that may be single-stranded or, more typically, double-stranded or a mixture of single- and double-stranded regions. In addition, the polynucleotide can be composed of triple-stranded regions comprising RNA or DNA or both RNA and DNA. A polynucleotide may also contain one or more modified bases or DNA or RNA backbones modified for stability or for other reasons. "Modified" bases include, for example, tritylated bases and unusual bases such as inosine. A variety of modifications can be made to DNA and RNA; thus, "polynucleotide" embraces chemically, enzymatically, or metabolically modified forms.

The polypeptide of the present invention can be composed of amino acids joined to each other by peptide bonds or modified peptide bonds, i.e., peptide isosteres, and may contain amino acids other than the 20 gene-encoded amino acids. The polypeptides may be modified by either natural processes, such as posttranslational processing, or by chemical modification techniques which are well known in the art. Such modifications are well described in basic texts and in more detailed monographs, as well as in a voluminous research literature. Modifications can occur anywhere in a polypeptide, including the peptide backbone, the amino acid side-chains and the amino or carboxyl termini. It will be appreciated that the same type of modification may be present in the same or varying degrees at several sites in a given polypeptide. Also, a given polypeptide may contain many types of modifications. Polypeptides may be branched, for example, as a result of ubiquitination, and

5

10

15

20

25

30

9

they may be cyclic, with or without branching. Cyclic, branched, and branched cyclic polypeptides may result from posttranslation natural processes or may be made by synthetic Modifications include acetylation, acylation, ADP-ribosylation, amidation, methods. covalent attachment of flavin, covalent attachment of a heme moiety, covalent attachment of a nucleotide or nucleotide derivative, covalent attachment of a lipid or lipid derivative, covalent attachment of phosphotidylinositol, cross-linking, cyclization, disulfide bond formation, demethylation, formation of covalent cross-links, formation of cysteine, formation of pyroglutamate, formylation, gamma-carboxylation, glycosylation, GPI anchor formation, hydroxylation, iodination, methylation, myristoylation, oxidation, pegylation, proteolytic processing, phosphorylation, prenylation, racemization, selenoylation, sulfation, transfer-RNA mediated addition of amino acids to proteins such as arginylation, and ubiquitination. (See, for instance, PROTEINS - STRUCTURE AND MOLECULAR PROPERTIES, 2nd Freeman and Company, New York (1993); Ed., T. E. Creighton, W. H. POSTTRANSLATIONAL COVALENT MODIFICATION OF PROTEINS, B. C. Johnson, Ed., Academic Press, New York, pgs. 1-12 (1983); Seifter et al., Meth Enzymol 182:626-646 (1990); Rattan et al., Ann NY Acad Sci 663:48-62 (1992).)

"SEQ ID NO:X" refers to a polynucleotide sequence while "SEQ ID NO:Y" refers to a polypeptide sequence. SEQ ID NO:X is identified by an integer specified in column 1 of Table 1. A translated open reading frame (ORF) encoded by polynucleotide SEQ ID NO:X, SEQ ID NO:Y, is shown in column nine of Table 1. There are 4277 colon and/or colon cancer related polynucleotide sequences described in Table 1 and shown in the sequence listing. Likewise there are 4277 colon and/or colon cancer related polypeptide sequences shown in the sequence listing, one polypeptide sequence for each of the polynucleotide sequences. The polynucleotide sequences are shown in the sequence listing immediately followed by all of the polypeptide sequences.

"A polypeptide having biological activity" refers to polypeptides exhibiting activity similar, but not necessarily identical to, an activity of a polypeptide of the present invention, including mature forms, as measured in a particular biological assay, with or without dose dependency. In the case where dose dependency does exist, it need not be identical to that of the polypeptide, but rather substantially similar to the dose-dependence in a given activity as compared to the polypeptide of the present invention (i.e., the candidate polypeptide will exhibit greater activity or not more than about 25-fold less and, preferably, not more than

5

10

15

20

about tenfold less activity, and most preferably, not more than about three-fold less activity relative to the polypeptide of the present invention).

PCT/US00/26524

The polynucleotides of the invention may be arrayed onto a nitrocellulose filter and screened with labelled mRNA which has been isolated from particular normal or diseased tissues, as described in Example 3. Known polynucleotide sequences are included in the array as hybridization controls, either because of their demonstrated tissue specificity or because they represent known surface molecules which may after further study show a predominant tissue expression and be useful antibody targets.

### Colon and/or Colon Cancer Related Polynucleotides and Polypeptides of the Invention

It has been discovered herein that the polynucleotides described in Table 1 are expressed at significantly enhanced levels in human colon and colon cancer tissues. Accordingly, such polynucleotides, polypeptides encoded by such polynucleotides, and antibodies specific for such polypeptides find use in the prediction, diagnosis, treatment, and prevention of disorders related to the colon, including, but not limited to colon cancer, as more fully described below.

Table 1 summarizes some of the polynucleotides encompassed by the invention (including polynucleotide sequences (SEQ ID NO:X) and the related cDNA clones (Clone ID:Z)) and further summarizes certain characteristics of these colon and/or colon cancer related polynucleotides, and the polypeptides encoded thereby.

Table 1

Seq ID No:X	Sequence/ Contig ID	Gene Name	Overlap	AA HGS SEQ ID Start No:Y	HGS Nucleotide Start End	cleotide	%I	% %	Clone ID:Z	Vector
-	390631	(AF051311) Ras-GAP SH3 binding protein [Homo sapiens] >sp O60606 O60606 RAS-GAP SH3 BINDING PROTEIN. Length = 449	gb AAC15705.	4278	8	326	96	96	HTWEP07	pSport1
2	410299			4279	75	251			HODBA26	Uni-ZAP XR
3	456200			4280	3	443			HPMEF95	HPMEF95 Uni-ZAP XR
4	456438	hypothetical protein (L1H 3" region) - human Length pir B34087 B3 = 1280	pir B34087 B3 4087	4281	-	513	43	58	HCFCY21	pSport1
5	467315			4282	278	412			HMKC008	pSport1
9	471563			4283	8	181			HBAGS04	pSport1
7	488131			4284	148	342			HALSQ75	Uni-ZAP XR
∞	490848			4285	241	522			HMVBD21	pSport1
6	969005	Similar to Volbox carteri extensin (S22697) [Homo sapiens] >gb AAD33052.1 AF134303_1 (AF134303) Scarl [Homo sapiens] >sp Q92558 Y269_HUMAN HYPOTHETICAL PROLINE-RICH PROTEIN KIAA0269. >sp AAD33052 AAD33052 Scarl. Length = 559	dbj BAA13399 .1	4286	2	187	78	78	HKIMD67	Lambda ZAP II
10	504559			4287	99	152			HOOAE34	pBluescript
11	506406			4288	34	420			HHSDD62	Uni-ZAP XR
12	506619			4289	85	1119			HSLGZ32	Uni-ZAP XR
13	507852			4290	211	456			HCENL15	Uni-ZAP XR
14	509423			4291	540	755			нсол138	HCQAI38 Lambda ZAP

HPMDT48 Uni-ZAP XR	pSport1	pBluescript SK-	Uni-ZAP XR	pBluescript SK-	HHSBA79 Uni-ZAP XR	HCQAQ89 Lambda ZAP	Uni-ZAP XR	Uni-ZAP XR	pBluescript SK-	HACCE33 Uni-ZAP XR	Uni-ZAP XR
HPMDT48	HADFX66		HEBBT54	H2CBG63	HHSBA79	НСОАО89	HELGJ91	HADBE91	HSAAX52	HACCE33	HE8DA85
										100	100
		_								100	100
316	345	434	819	232		388	363	239	450	403	1626
197	172	318	808	11	159	224	241	3	250	2	562
4292	4293	4294	4295	4296	4297	4298	4299	4300	4301	4302	4303
										dbj BAA00931 .1	emb CAA4291 6.1
										long-chain acyl-CoA synthetase [Homo sapiens] >pir[JX0202]JX0202 long-chain-fatty-acidCoA ligase (EC 6.2.1.3) - human >sp P33121 LCFB_HUMAN LONG-CHAIN- FATTY-ACIDCOA LIGASE 2 (EC 6.2.1.3) (LONG-CHAIN ACYL-COA SYNTHETASE 2) (LACS 2). Length = 698	5-aminolevulinate synthase precursor [Homo sapiens] >emb CAA15886.1  (AL020991) d1884M20.2 [Homo sapiens] >gb AAC39838.1  (AF068624) 5-aminolevulinate synthase 2 [Homo sapiens] >pir S16347 SYHUAE 5-aminolevulinate synthase (EC 2.3.1.37) precursor, erythro
509734	958605	524721	524901	527600	527827	529050	529465	530612	530773	532810	533242
15	16	17	18	19	20	21	22	23	24	25	26

pBluescript	pSport1	Lambda ZAP II	Uni-ZAP XR	ZAP Express	Uni-ZAP XR	pCMVSport 2.0	pSport1
HSKII86	HUSGI39	HKIMB44	HBMVJ62	HBXFC78	HE2FR32	HKACD58	HDAAB62
93		100				81	83
92		100				08	77
622	285	240	611	390	262	1208	339
140	259	133	345	184	41	969	208
4304	4305	4306	4307	4308	4309	4310	4311
gb AAA35871.		gb AAA35567. 1				emb CAB4324 7.1	emb CAA3443
guanylate binding protein isoform I [Homo sapiens] >pir A41268 A41268 guanine nucleotide-binding protein 1 - human >sp P32455 GBP1_HUMAN INTERFERON-INDUCED GUANYLATE-BINDING PROTEIN 1 (GUANINE NUCLEOTIDE-BINDING PROTEIN 1). Length = 592		aspartyl-tRNA synthetase [Homo sapiens] >pir A34393 SYHUDT aspartatetRNA ligase (EC 6.1.1.12) - human >sp P14868 SYD_HUMAN ASPARTYL-TRNA SYNTHETASE (EC 6.1.1.12) (ASPARTATETRNA LIGASE) (ASPRS). Length = 500				(AL050037) hypothetical protein [Homo sapiens] >emb[CAB43247.1  (AL050037) hypothetical protein [Homo sapiens] >pir[T08715]T08715 hypothetical protein DKFZp566I1024.1 - human (fragment) >sp[Q9Y405 Q9Y405 HYPOTHETICAL 34.8 KD PROTEIN (FRAGMENT). Length = 3	precursor polypeptide (AA -29 to 315) [Homo sapiens] >pir S14902 DEHUMT methylenetetrahydrofolate dehydrogenase (NAD+) (EC 1.5.1.15) / methenyltetrahydrofolate cyclohydrolase (EC 3.5.4.9) precursor - human >sp P 13995 MTDC_HUMAN BIFUNCTIONAL METHYLENETETRA
541126	542268	547920	549642	550207	552115	552465	554369
27	28	29	30	31	32	33	34

HEPBA24 Uni-ZAP XR	pCMVSport 2.0	pBluescript SK-	pBluescript SK-	HCQAT53 Lambda ZAP	HETDN09 Uni-ZAP XR	pBluescript SK-	pCMVSport 3.0	pBluescript SK-	pCMVSport 3.0
HEPBA24	HOGBL08	HCYBD62	H2CBD20	HCQAT53	HETDN09	HCYBE04	HDPFI14	НЈВСД90	HAJAB40
		62	63		91		96	89	9
		62	51		91	_	96	41	48
009	530	379	688	254	628	423	892	423	226
379	222	2	773	т	2	283	2	31	2
4312	4313	4314	4315	4316	4317	4318	4319	4320	4321
		emb CAA6176 1.1	gb AAA88038. 1		gb AAA35899. 1		dbj BAA21367 .1	emb CAA9699 3.1	emb CAB1168 0.1
			protein [Homo sapiens] >splQ14288 Q14288 HYPOTHETICAL PROTEIN (FRAGMENT). Length = 641		gamma-glutamyl transpeptidase [Homo sapiens] >emb CAA07485.1  (AJ007378) gamma-glutamyltransferase [Homo sapiens] {SUB 193-244} >emb CAA07487.1  (AJ007380) gamma-glutamyltransferase [Homo sapiens] {SUB 296-340} Length = 569		NPAT [Homo sapiens] >dbj BAA11861.1  NPAT [Homo sapiens] >sp Q16580 Q16580 NPAT (E14 AND A-T PROTEINS). Length = 1427	ORF YGR010w [Saccharomyces cerevisiae] >pir S64299 S64299 probable membrane protein YGR010w - yeast (Saccharomyces cerevisiae) >sp P53204 YG15_YEAST HYPOTHETICAL 44.9 KD PROTEIN IN SEC9-MSB2 INTERGENIC REGION. Length = 395	methionyl-trna synthetase, mitochondrial [Schizosaccharomyces pombe] >pir[T38454 T38454 methionyl-trna synthetase, mitochondrial - fission yeast (Schizosaccharomyces pombe) >sp[O14000 O14000 PUTATIVE METHIONYL-TRNA SYNTHETASE (EC 6.1.1.10) (METHIONINE-TR
557152	557230	558366	570796	573181	573199	573793	573796	574094	574927
35	36	37	38	39	40	41	42	43	44

pBluescript SK-	pCMVSport 3.0	HLMMR55 Lambda ZAP	HNFGN91 Uni-ZAP XR	pSport1	Lambda ZAP II	Uni-ZAP XR	HNHDV16 Uni-ZAP XR	Uni-ZAP XR	HSAVM80 Uni-ZAP XR	pSport1	Uni-ZAP XR	HELHD03 Uni-ZAP XR	pBluescript	pSport1	Uni-ZAP XR
H2MCA74	HWBAX42	HLMMR55	HNFGN91	HTWD190	HCQAB18	HELHI45	HNHDV16	HOAAD32	HSAVM80	HWLMAS1	не8во01	НЕГНД03	HBMCT70	HLYDF04	HDSAP04
	81						08			98					
	08						64			88					
429	401	255	322	557	256	239	406	361	285	408	231	428	340	157	321
151	3	1	59	345	83	3	528	2	10	_	55	273	215	5	136
4322	4323	4324	4325	4326	4327	4328	4329	4330	4331	4332	4333	4334	4335	4336	4337
	gb AAB60340. 1						gb AAA03341. 1			emb CAA8054					
	ATP:citrate lyase [Homo sapiens] >sp Q13037 Q13037 ATP:CITRATE LYASE. Length = 1101						ribosomal protein L23a [Homo sapiens] gb AAA03341. >gb AAA35681.1  homology to rat ribosomal protein 1  L23 [Homo sapiens] {SUB 10-156} Length = 156			M130 antigen [Homo sapiens] >emb CAB45233.1  CD163 [Homo sapiens] >pir 138003 S36077 M130 antigen - human >sp Q07898 Q07898 M130 ANTIGEN PRECURSOR. Length = 1116					
575139	575591	576132	577390	577685	578079	578660	280860	581143	584899	699009	611839	614078	614554	615029	615590
45	46	47	48	49	20	51	52	53	54	55	56	57	58	59	09

pCMVSport 3.0	HCQBH72 Lambda ZAP	Uni-ZAP XR	HNHEU34 Uni-ZAP XR	pCMVSport 3.0	pCMVSport 3.0	Lambda ZAP II	pSport1	n	HCQAW11 Lambda ZAP II	Uni-ZAP XR
HWBFZ21	нсовн72	HELGH31	HNHEU34	HJMAF30	нwввк93	HFXAK32	HUSIT18	HMWBH51	HCQAW11	HPRAS01
85			89		63	73		78	100	
83			89		49	<i>L</i> 9		78	100	
423	764	604	573	254	821	1400	1040	308	198	153
-	492	401	199	105	£	1185	759	33	_	-
4338	4339	4340	4341	4342	4343	4344	4345	4346	4347	4348
gb AAC68903.			gb AAD45830. 1 AC0048		gb AAC15584. 1	gb AAF22026. 1 AF1180		gb AAB41942. 1	emb CAB4328	
(AF098799) RanBP7/importin 7 [Homo sapiens] >sp[O95373 O95373 RANBP7/IMPORTIN 7. >emb[CAB70698.1] (AL137335) hypothetical protein [Homo sapiens] {SUB 831-1038} Length = 1038			(AC004876) similar to neuro-endocrine specific protein VGF; similar to CAA73210 (PID:g2244659) [Homo sapiens] >sp AAD45830 AAD45830 WUGSC:H_DJ0747G18.3 protein. Length = 615		(AF026198) putative protein 2 [Fugu rubripes] >pir[T30536]T30536 hypothetical protein 2 - Fugu rubripes (fragment) >sp 073698 073698 HYPOTHETICAL 21.5 KD PROTEIN (FRAGMENT). Length = 187	(AF118082) PRO1902 [Homo sapiens] >sp AAF22026 AAF22026 PRO1902. Length = 84		K-ras oncogene protein [Homo sapiens] Length = 188	(AL050120) hypothetical protein [Homo sapiens] >emb CAB43281.1  (AL050120) hypothetical protein [Homo sapiens] >pir T08766 T08766 hypothetical protein DKFZp586D211.1 - human (fragment) >sp CAB43281 CAB43281 Hypothetical 15.3 kd protein (fragment). Length	
630230	637548	637605	638125	638188	638249	638319	651380	651876	653175	655544
61	62	63	49	65	99	29	89	69	70	71

pCMVSport 3.0	pCMVSport 3.0	pCMVSport 3.0	Jni-ZAP XR	Uni-ZAP XR	Uni-ZAP XR	Uni-ZAP XR	ambda ZAP II	pCMVSport 3.0	PCRII	pCMVSport 2.0	Cambda ZAP II
HWBBC13	HNTBM67	HDPKC15	HMAHP16 Uni-ZAP XR	HCE1D45	HBIBV81	HSXBP02	HCQCO19 Lambda ZAP	HHENT19	100 HMTMB52	HOGAL19	HCQAG50 Lambda ZAP
	94			84					100		69
	94			84					100		59
537	1058	491	341	279	448	277	826	342	535	171	280
331	120	321	132		239	2	407	169	2	92	110
4349	4350	4351	4352	4353	4354	4355	4356	4357	4358	4359	4360
	dbj BAA77295			gb AAF01349. 1 AC0050					gb AAD34132. 1 AF1518		dbj BAA91271 .1
	(AB014509) Nck-associated protein 1 (Nap1) [Homo dbj BAA77295 sapiens] >sp Q9Y2A7 Q9Y2A7 NCK-ASSOCIATED PROTEIN 1 (NAP1). Length = 1128			(AC005003) similar to zinc finger protein MAZ [Homo sapiens]; similar to AAB04121.1 (PID:g995935) >sp AAF01349 AAF01349 WUGSC:H_DJ400N23.1 protein. >emb CAB51404.1  (AL096880) hypothetical protein [Homo sapiens] {SUB 26-641} Length = 641					(AF151895) CGI-137 protein [Homo sapiens] >gb AAF14860.1 AF110777_1 (AF110777) adrenal gland protein AD-004 [Homo sapiens] >sp Q9Y3D8 YCD7_HUMAN HYPOTHETICAL PROTEIN CGI-137. >sp AAF14860 AAF14860 Adrenal gland protein AD-004. Length = 172		(AK000585) unnamed protein product [Homo sapiens] Length = 285
656722	659801	660020	661600	664481	665154	062999	668040	985899	668717	668753	671361
72	73	74	75	76	77	78	79	08	81	82	83

HDPLC22 pCMVSport 3.0	HBMX090 Uni-ZAP XR	Lambda ZAP II	HE8AG73 Uni-ZAP XR	pBluescript SK-	pCMVSport 2.0	ZAP Express	pSport1
HDPLC22	HBMXO90	HLMIS22	HE8AG73	HCYBF14	HKAAS37	HBXFP72	HFIYP15
	94		86			81	51
	94		86			72	38
405	1243	379	675	404	168	1167	978
280	71	194	-	63	-	-	394
4361	4362	4363	4364	4365	4366	4367	4368
	dbj BAA25452 .1		gb AAD30288. 1 AF1364			emb CAB5926 1.1	emb CAB5368
	(AB011098) KIAA0526 protein [Homo sapiens] >emb CAA69942.1  serine palmitoyltransferase, subunit II [Homo sapiens] >gb AAD09621.1  (AF111168) serine palmitoyl transferase, subunit II [Homo sapiens] >sp O15270 LCB2_HUMAN SERINE PALMITOYLTRANSFERASE 2 (EC 2		(AF136450) goodpasture antigen-binding protein [Homo sapiens] >splQ9Y5P4 Q9Y5P4 GOODPASTURE ANTIGEN-BINDING PROTEIN (EC 2.7.1.37). Length = 624			(AL122091) hypothetical protein [Homo sapiens] >emb CAB59261.1  (AL122091) hypothetical protein [Homo sapiens] >pir T34522 T34522 hypothetical protein DKFZp566D244.1 - human (fragment) >sp CAB59261 CAB59261 Hypothetical 64.0 kd protein (fragment). Length	(AL110226) hypothetical protein [Homo sapiens] >emb CAB53684.1  (AL110226) hypothetical protein [Homo sapiens] >pir T14764 T14764 hypothetical protein DKFZp434H204.1 - human (fragment) >sp CAB53684 CAB53684 Hypothetical 96.7 kd protein (fragment). Length
674203	674745	674761	677212	683259	685895	688040	688044
84	\$	98	87	88	68	06	91

HEBAG86 Uni-ZAP XR	pCMVSport 3.0	pCMVSport 3.0	HE2OC31 Uni-ZAP XR	HTXKQ20 Uni-ZAP XR	HE2OK20 Uni-ZAP XR	HMWIW31 Uni-ZAP XR	HCEEH33 Uni-ZAP XR	HAGBL85 Uni-ZAP XR	pCMVSport 3.0	pBluescript SK-	Uni-ZAP XR	Uni-ZAP XR	ZAP Express	Uni-ZAP XR	Uni-ZAP XR
	HLDNM81	HARNC71	HE20C31	нтхк020	HE20K20	HMWIW31	нсеен33	HAGBL85	HLWAY38	H2LAN34	HBMXT67	HE2IE28	HBXCG73	HATAN68	HAGDD59
100				62					96	74			68	86	
100				41					96	64			88	97	
276	418	411	428	191	135	242	391	258	685	211	1267	144	703	412	159
	230	139	237	3	34	3	113	-	2	2	914	9/	185	2	1
4369	4370	4371	4372	4373	4374	4375	4376	4377	4378	4379	4380	4381	4382	4383	4384
gb AAC39891.				pir JC4296 JC4 296					gb AAD38244. . 1 AC0077	emb CAB4218 7.1			gb AAC39757.	gb AAC00205.	
(AF047440) ribosomal protein L33-like protein [Homo sapiens] >sp O75394 O75394 RIBOSOMAL PROTEIN L33-LIKE PROTEIN. Length = 65				ring finger protein - fruit fly (Drosophila melanogaster) Length = 222					(AC007785) BC282485_1 [Homo sapiens] >sp Q9Y6R9 Q9Y6R9 BC282485_1 (FRAGMENT). Length = 477	unnamed protein product [unidentified] Length = 309 emb CAB4218 7.1			spectrin SH3 domain binding protein I [Homo sapiens] >sp[O76049 O76049 SPECTRIN SH3 DOMAIN BINDING PROTEIN 1. Length = 508	PRAJA1 [Mus musculus] >sp O55176 O55176 PRAJA1. Length = 424	
688077	691124	691721	693582	200969	697955	890869	702853	703700	705461	705692	706204	707161	707464	709015	709518
92	93	94	95	96	26	86	66	100	101	102	103	104	105	106	107

Uni-ZAP XR	HSNAL84 Uni-ZAP XR	pSport1	pBluescript	Uni-ZAP XR	Uni-ZAP XR	pSport1	HFXHM92 Lambda ZAP	HHSGE44 Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1
HBJF165	HSNAL84	HCRND41	HPXAA41	HHSFO42	HCEIE94	HWLQA43	HFXHM92	HHSGE44	HWLQ133	HFIAW90	HOSEP43	HUSGY48	HSLEC18	HUFAC36
		85		11					37				61	55
		79		75					37				42	37
323	427	189	574	149	992	168	241	703	402	299	190	101	1319	1547
102	182	-	254	<u>س</u>	009	10	29	299	-	527	71	3	En .	1236
4385	4386	4387	4388	4389	4390	4391	4392	4393	4394	4395	4396	4397	4398	4399
		sp P39194 AL U7_HUMAN		sp P39189 AL U2_HUMAN					dbj BAA91415 .1				dbj BAA13673 .1	gb AAA49527. 1
		IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593		IIII ALU SUBFAMILY SB WARNING ENTRY IIII sp P39189 AL Length = 587					(AK000900) unnamed protein product [Homo sapiens] Length = 136				cerebroside sulfotransferase [Homo sapiens] >dbj BAA89503.1  (AB029901) cerebroside sulfotransferase [Homo sapiens] >gb AAD50517.1 AC005006_2 (AC005006) cerebroside sulfotransferase [Homo sapiens] >sp Q9999 Q9999 CEREBROSIDE SULFOTRANSFERASE. >sp BAA895	olfactomedin [Rana catesbeiana] >pir A47442 A47442 olfactomedin precursor - bullfrog >sp Q07081 OLFM_RANCA OLFACTOMEDIN PRECURSOR (OLFACTORY MUCUS PROTEIN). Length = 464
711769	711840	711878	712638	713301	714156	714877	715343	716212	717222	718259	719829	721985	722249	722258
801	109	110	111	112	113	114	115	116	117	118	119	120	121	122

	put. ting protein [fromo sapiens] >sp Q99579 Q99579 PUTATIVE RING PROTEIN. Length = 236	emb CAA6916 5.1	I	571	143	83	87	HHFHB49	Ď
725110			4401	174	323			HFIBH05	pSport1
725201	(AB020676) KIAA0869 protein [Homo sapiens] >sp O94946 O94946 KIAA0869 PROTEIN (FRAGMENT). Length = 888	dbj BAA74892 .1	4402		294	86	100	HKIAA57	Uni-ZAP XR
726122			4403	309	530			HRKAB52	pBluescript
727365			4404	1	915			HPCAN95	Uni-ZAP XR
729143			4405	609	821			HCQCV54	Lambda ZAP II
729231			4406	345	602			HLJEA54	pCMVSport 1
731881	IIII ALU SUBFAMILY J WARNING ENTRY IIII Length = 591	sp P39188 AL U1_HUMAN	4407	361	462	56	92	HTWCR70	pSport1
732280	(AB002349) KIAA0351 [Homo sapiens] >sp 015059 015059 KIAA0351. Length = 557	dbj BAA20808	4408	155	298	81	82	HSXDD55	Uni-ZAP XR
732932			4409	295	483			HSTAB63	Uni-ZAP XR
733034	expressed-Xq28STS protein [Homo sapiens] Length = 358	gb AAF33529. 1 U82695	4410	305	553	98	68	H6BSI11	Uni-ZAP XR
734012			4411	49	180			НДОРР57	pCMVSport 3.0
735603			4412	991	1224			HAGEX59	HAGEX59 Uni-ZAP XR
739061			4413	436	621			HAVMG19	Other
741134	protein [Homo sapiens] >sp Q14288 Q14288 HYPOTHETICAL PROTEIN (FRAGMENT). Length = 641	gb AAA88038. 1	4414	929	314	19	69	HLEAL50	Uni-ZAP XR
741257	protein [Homo sapiens] >sp Q14287 Q14287 HYPOTHETICAL PROTEIN (FRAGMENT). Length = 157	gb AAA88036. 1	4415	136	195	41	52	HCPAC07	Uni-ZAP XR
741804			4416	196	441			HOSEQ61	Uni-ZAP XR

	742220	initiation factor 5A [Gallus gallus] >pir [50227 A42156 translation initiation factor eIF- 5A 1 - chicken >sp Q07460 IF51_CHICK INITIATION FACTOR 5A-1 (EIF-5A) (EIF-4D). Length = 153	gb AAA17444.   1	4417	<i>د</i>	455	83	88	HCROB09	pSport1
	744605			4418	255	479			HFIZP62	pSport1
	744687			4419	231	374			HBMTK19	HBMTK19 Uni-ZAP XR
	745368			4420	_	114			HAGDG84	HAGDG84 Uni-ZAP XR
i	747870			4421	343	519			HCABQ86	Uni-ZAP XR
t .	750486			4422	694	268			HSAXE65	Uni-ZAP XR
t .	751119	Impact [Mus musculus] >sp O55091 O55091 IMPACT PROTEIN. Length = 318	dbj BAA35139  .1	4423	2	481	89	77	HE80C67	Uni-ZAP XR
•	752557			4424	1	210			НКАНА68	pCMVSport 2.0
	753226	PROTEIN (FRAGMENT). Length = 184	sp Q29229 Q2 9229	4425	25	810	38	51	HSFAG23	Uni-ZAP XR
	754269			4426	737	949			HDTAT69	pCMVSport 2.0
	756466	(AF099731) connexin 31.1 [Homo sapiens] >sp O95377 CXB5_HUMAN GAP JUNCTION BETA-5 PROTEIN (CONNEXIN 31.1) (CX31.1). Length = 273	gb AAC95472. 1	4427	3	293	96	96	HAICM70	HAICM70 Uni-ZAP XR
	756538	homologous to mouse Rsu-1; putative [Homo sapiens] >pir I60122 I60122 rsu-1 homolog - human >sp Q15404 RSU1_HUMAN RAS SUPPRESSOR PROTEIN 1 (RSU-1) (RSP-1 PROTEIN) (RSP-1). Length = 277	gb AAA60292. 1	4428	93	1151	100	100	HMCGF70	HMCGF70 Uni-ZAP XR
	756649	The ha2022 gene product is novel. [Homo sapiens] >sp Q14699 Y084_HUMAN HYPOTHETICAL PROTEIN KIAA0084 (HA2022) (FRAGMENT). Length = 648	dbj BAA07644 .1	4429		135	100	100	HE8EX74	Uni-ZAP XR

HACBN11 Uni-ZAP XR	HTTBS70 Uni-ZAP XR	4 pSport1	Uni-ZAP XR	pCMVSport 3.0	HRABS72 pCMVSport 3.0	HYAAX74 pCMVSport 3.0	9 pBluescript	5 pBluescript	5 pBluescript SK-	6 Uni-ZAP XR	HEIBB38 Uni-ZAP XR	6 pCMVSport 3.0	7 Uni-ZAP XR	HTLDW36 Uni-ZAP XR
	HTTBS70	HCRNF04	HETIS94	HDPXJ71	HRABS72	HYAAX7	HSKXC19	HF6SG75	HCYBG95	HCECT76	HEIBB38	HHEMK76	HE9PB77	HTLDW3
85	68			79								66		100
84	87			79								86		100
362	734	385	622	763	377	183	661	243	472	497	919	465	664	611
m	e	86	365	2	153	43	2	112	332	327	632	-	374	129
4430	4431	4432	4433	4434	4435	4436	4437	4438	4439	4440	4441	4442	4443	4444
dbj BAA91704 .1	emb CAA7084 4.1			dbj BAA91356 .1								emb CAA6691 1.1		gb AAA90924.  1
(AK001459) unnamed protein product [Homo sapiens] Length = 245	sapiens] >sp O95664 O95664 igth = 758			(AK000743) unnamed protein product [Homo sapiens] Length = 573								M-phase phosphoprotein 9 [Homo sapiens] >sp Q99550 MPP9_HUMAN M-PHASE PHOSPHOPROTEIN 9 (FRAGMENT). Length = 214		DNA polymerase epsilon catalytic subunit [Homo sapiens] >pir G02434 G02434 DNA-directed DNA polymerase (EC 2.7.7.7) epsilon catalytic chain -
757213	757508	757532	757980	760141	761491	761724	762027	764179	766961	767593	768034	769965	771486	772044
153	154	155	156	157	158	159	160	191	162	163	164	165	166	167

772357	357		gb AAA37536. 1	4445	403	738	98	93	HMWHN43	93 HMWHN43 Uni-ZAP XR
		>splQ0/139 EC12_MOUSE_EC12 PROTEIN (ECT2 ONCOGENE). Length = 738								
772876	928	(AK000771) unnamed protein product [Homo sapiens] Length = 202	dbj BAA91373 .1	4446	59	820	66	66	HUSIR49	pSport1
774019	610	apiens] >sp 095792 095792 .1 KD PROTEIN. Length =	gb AAD17528.	4447	14	520	75	75	НЕ9НҮ44	Uni-ZAP XR
774244	244	(AL009196) /prediction=(method:"genefinder", version:"084"); /prediction=(method:"genscan", version:"1.0"); /match=(desc:"LD09991.5prime LD Drosophila melanogaster embryo BlueScript Drosophila melanogaster cDNA clone LD09991 5prime, mRNA seque>>	emb CAA1571 2.1	4448	2	1123	71	98	HTTEL19	Uni-ZAP XR
774516	516	(AK000482) unnamed protein product [Homo sapiens] Length = 572	dbj BAA91194 .1	4449	1	438	54	74	HMCFS02	HMCFS02 Uni-ZAP XR
775355	355			4450	1599	1781			HDTBY31	pCMVSport 2.0
775367	367	:		4451	142	228			HUSXP15	pSport1
175791	791			4452	588	911			HSAWS31	Uni-ZAP XR
777319	319			4453	283	477			HE8OV83	HE8OV83 Uni-ZAP XR
778434	434	stress-activated protein kinase-3 [Homo sapiens] >emb CAB51538.1  (AL022328) dJ402G11.1 (mitogen activated protein kinase 12 (PRKM11)) [Homo sapiens] >sp P53778 MK12_HUMAN MITOGEN-ACTIVATED PROTEIN KINASE 12 (EC 2.7.1) (EXTRACELLULAR SIGNAL- REGULATED KI	emb CAA7151	4454	2	880	96	96	HL3AD81	HL3AD81 Uni-ZAP XR

178	778583	(AF000198) weak similarity to HSP90 [Caenorhabditis elegans] >pir[T15138[T15138] hypothetical protein T28F2.4 - Caenorhabditis elegans >sp O01658 O01658 SIMILARITY TO HSP90. Length = 817	gb AAB53055.	4455	т	248	29	52	<b>ННЕ</b> RQ03	pCMVSport
621	779480			4456	331	546			HTXFI40	Uni-ZAP XR
180	779588			4457	49	162			HBIMB82	pCMVSport 3.0
181	781085			4458	2300	2641			HTTEW79	HTTEW79 Uni-ZAP XR
182	781286			4459	149	289			HLJB183	pCMVSport 1
183	781366			4460	261	909			HSAWU83	Uni-ZAP XR
184	781376			4461	23	340			HADFW62	pSport1
185	781832			4462	587	733			HSNAK79	Uni-ZAP XR
186	782276	(AB032969) KIAA1143 protein [Homo sapiens] >sp BAA86457 BAA86457 KIAA1143 protein (fragment). Length = 116	dbj BAA86457	4463	2	472	17	71	HSUBX87	Uni-ZAP XR
187	782358			4464	1008	1289			HATEF13	Uni-ZAP XR
188	783413	D9 splice variant 1 [Mus musculus] >sp 008693 008693 D9 SPLICE VARIANT 1. Length = 111	gb AAB53635. 1	4465	-	591	08	88	HEBFR23	Uni-ZAP XR
189	783668			4466	171	464			HARMP12	pCMVSport 3.0
190	783677			4467	240	380			HJMBT13	pCMVSport 3.0
161	785087	beta-galactosidase alpha peptide [Cloning vector pSport2] Length = 114	gb AAA67217.	4468	909	815	95	95	HEAAK74	HEAAK74 Uni-ZAP XR
192	785328	unnamed protein product [unidentified] >emb CAB42218.1  unnamed protein product [unidentified] {SUB 62-446} Length = 446	emb CAB4221 2.1	4469	388	609	91	95	HAMGI86	pCMVSport 3.0

	•													
HDPCN86 pCMVSport 3.0	HMCGH90 Uni-ZAP XR	pCMVSport 1	Uni-ZAP XR	pSport1	pCMVSport 1	pSport1	Uni-ZAP XR	pBluescript SK-	HE9SD26 Uni-ZAP XR	pSport1	pCMVSport 3.0	pSport1	pCMVSport 3.0	pSport1
HDPCN86	<b>НМС</b> БН90	ННВҒМЗЗ	HSLF109	HFIAX76	HLICN93	HCFBE51	HFEAU63	HAFBC92	HE9SD26	HFIZG43	HDPUX67	HVAAA93	HAMFQ15	HADCW71
	86	52					94	98					93	78
	86	30					06	96					92	89
535	79	388	209	410	320	314	1989	313	545	2102	283	222	470	465
2	399	2	108	141	192	3	2249	2	168	1893	161	64	231	1
4470	4471	4472	4473	4474	4475	4476	4477	4478	4479	4480	4481	4482	4483	4484
	dbj BAA78730 .1	emb CAB6263 1.1					emb CAA7130 9.1	dbj BAA11319 .1					dbj BAA21720 .1	gb AAB49620. 1
	(AB028639) PalBH [Homo sapiens] >sp Q9Y6W3 Q9Y6W3 PALBH (EC 3.4.22.17). Length = 813	outative protein [Arabidopsis thaliana] 11 CAB62631 Hypothetical 29.5 kd the = 263						PAP-1 [Mus musculus] >sp P97762 P97762 PAP-1. Length = 213					factor Length =	PACT [Mus musculus] >sp P70287 P70287 RETINOBLASTOMA BINDING PROTEIN 6 (PACT) (FRAGMENT). Length = 1587
785465	788626	788838	789286	789419	789631	789872	790190	790547	791155	791220	791749	792034	792557	792624
193	194	195	196	197	198	199	200	201	202	203	204	205	206	207

pSport1	HLQAX49 Lambda ZAP	HMAJP26 Uni-ZAP XR	Uni-ZAP XR	pSport1	Uni-ZAP XR	pBluescript SK-	pSport1	pCMVSport 3.0	pBluescript	Uni-ZAP XR
HCHMB04	HLQAX49	HMAJP26	HBJEA52	HPSNE17	нтесв93	HCYBF25	HGAMA30	HRACH60	HNFIZ54	HMSCL38
57		100			79	06	93		75	64
47		96			75	85	93		73	55
3	323	101	1399	699	174	1450	1655	251	273	2899
215	162	3	842	268	-	1376	603	3	-	2792
4485	4486	4487	4488	4489	4490	4491	4492	4493	4494	4495
emb CAA4164 0.1		gb AAA20069.			sp P39193 AL U6_HUMAN	sp P39189 AL U2_HUMAN	dbj BAA22165 .1		gb AAA36388.	dbj BAA91205 .1
actVA 4 [Streptomyces coelicolor A3(2)] >pir S18542 S18542 hypothetical protein 4 - Streptomyces coelicolor >sp Q53906 Q53906 ACTVA REGION GENES OF THE ACTINORHODIN BIOSYNTHETIC GENE CLUSTER. Length = 294		cytochrome c oxidase subunit II [Pan troglodytes] >sp P26457 COX2_PANPA CYTOCHROME COXIDASE POLYPEPTIDE II (EC 1.9.3.1). Length = 227			!!!! ALU SUBFAMILY SP WARNING ENTRY !!!! sp P39193 AL Length = 593	IIII ALU SUBFAMILY SB WARNING ENTRY IIII   sp P39189 AL   Length = 587	(AB006781) galectin-4 [Homo sapiens] >gb AAB86590.1  galectin-4 [Homo sapiens] >gb AAC51763.1  (AF014838) galectin-4 [Homo sapiens] >sp P56470 LEG4_HUMAN GALECTIN-4 (LACTOSE-BINDING LECTIN 4) (L-36 LACTOSE BINDING PROTEIN) (L36LBP). >sp AAB86590 AAB86590		oncostatin M [Homo sapiens] >gb AAC05173.1  (AC004264) oncostatin M precursor [Homo sapiens] >pir A32489 A32489 oncostatin M precursor - human >sp P13725 ONCM_HUMAN ONCOSTATIN M PRECURSOR (OSM). Length = 252	(AK000496) unnamed protein product [Homo sapiens] Length = 239
793437	795184	795744	796023	796181	797079	797477	797486	797747	800085	801919
208	509	210	211	212	213	214	215	216	217	218

t			<u>a</u>	8	<u> </u>	8	Ħ	8
pCMVSport 3.0	pSport1	pSport1	HMEBY61 Lambda ZAP	HETDK50 Uni-ZAP XR	Uni-ZAP XR	HLTDL01 Uni-ZAP XR	pCMVSport 2.0	Uni-ZAP XR
pCN	P.S		Lam	Uni	Uni-	Uni	pCN	
HDQGA42	HFIIY89	НВОЕВ83	BY61	DK50	нѕієн63	DL01	HKAJJ29	HTPCH84
НДО	HEI	HBO	HME	HET	HSI	HLT	HK/	HTP
		06	70	63	78		100	98
		06	47	43	09		100	98
200	349	1333	1257	1339	541	87	864	1023
51	125	2	184	2	2	1	151	
4496	4497	4498	4499	4500	4501	4502	4503	4504
		7963	5089.	4419	7358.		1162	3056.
		emb CAA7963 5.1	gb AAC45089. 1	dbj BAA24419 .1¦	gb AAC77358. 1		dbj BAA91162 .1	gb AAD03056. 1
		em - 5.1	dg				.db <u>⊢.</u>	
		thrombospondin-4 [Homo sapiens] emb >pir A55710 TSHUP4 thrombospondin 4 precursor - 5.1  human >sp P35443 TSP4_HUMAN THROMBOSPONDIN 4 PRECURSOR. Length = 961	VCP-like ATPase [Thermoplasma acidophilum]   >pir T37458 T37458 VCP-like ATPase - Thermoplasma acidophilum >sp O05209 O05209   VCP-LIKE ATPASE. Length = 745	(AB000549) alpha,-antitrypsin-like protein [Tamias sibricus] >sp O54760 ALSI_TAMSI ALPHA-1-ANTITRYPSIN-LIKE PROTEIN CM55-SI PRECURSOR. Length = 413	(AF007791) secreted cement gland protein XAG-2 homolog [Homo sapiens] >gb AAC82614.1  (AF038451) secreted cement gland protein XAG-2 homolog [Homo sapiens] >gb AAF22484.1 AF088867_1 (AF088867) putative secreted protein XAG [Homo sapiens] >pir JE0350 JE035		(AK000432) unnamed protein product [Homo sapiens] Length = 379	(AF104419) decoy receptor 3 [Homo sapiens] >gb AAD29688.1 AF134240_1 (AF134240) tumor necrosis factor receptor homolog [Homo sapiens] >gb AAF33685.1 AF217793_1 (AF217793) M68C [Homo sapiens] >gb AAF33686.1 AF217794_1 (AF217794) M68E [Homo sapiens] >gb AAF
805448	069908	810870	811047	812745	812755	812871	813482	815696
219	220	221	222	223	224	225	226	227

pCMVSport 3.0	pCMVSport 3.0	Uni-ZAP XR	pSport1	HAIDQ59 Uni-ZAP XR	pCMVSport 2.0	HBNAP17 Uni-ZAP XR	pSport1	Uni-ZAP XR	Uni-ZAP XR	pSport1
HWDAC26 pCMVSport 3.0	HMUB122	HMSDI67	HWLEZ80	НАІDQ59	HTJNI76	HBNAP17	HWLFM26	HPWBE34	HPICC36	HFOYL30
76		58	81	88			83	86		100
57		44	63	85			83	95		66
1323	1594	1202	964	889	411	838	883	<i>L</i>	350	661
1057	1370	1279	764	2	190	716	20	390	222	2
4505	4506	4507	4508	4509	4510	4511	4512	4513	4514	4515
gb AAA88036.		gb AAB02291. 1	dbj BAA13251 .1	emb CAB6982 7.1	:		gb AAF19050. 1	dbj BAA14834 .1		emb CAA6729 5.1
protein [Homo sapiens] >sp[Q14287]Q14287 HYPOTHETICAL PROTEIN (FRAGMENT). Length = 157		reverse transcriptase [Homo sapiens] Length = 361	GS3786 [Homo sapiens] >gb AAD54511.1 AC006364_1 (AC006364) GS3786 [Homo sapiens] >sp Q92520 G786_HUMAN PROTEIN GS3786. >sp AAD54511 AAD54511 GS3786. Length = 227	(AJ271442) Tspan-2 protein [Rattus norvegicus] >sp CAB69827 CAB69827 Tspan-2 protein. Length = 221			(AF067797) aquaporin 8 [Homo sapiens] >sp AAF19050 AAF19050 Aquaporin 8. Length = 261	ORF_ID:o255#5; similar to [SwissProt Accession Number P45576] [Escherichia coli] >gb AAC74362.1  (AE000226) putative heat shock protein [Escherichia coli] >pir C64876 C64876 yciM protein precursor - Escherichia coli >sp P45576 YCIM_ECOLI HYPOTHETICAL 44.5		RNA helicase [Homo sapiens] >pir S71758 S71758 DEAD box protein MrDb, Myc-regulated - human >sp Q92732 Q92732 RNA HELICASE. Length = 610
821335	824071	827298	827315	827562	827721	827740	828180	828552	828670	828919
228	229	230	231	232	233	234	235	236	237	238

pSport1	Uni-ZAP XR	pSporti	Uni-ZAP XR	HYAAS90 pCMVSport 3.0	HLDCP20 pCMVSport 3.0	pSport1	pSport1	pSport1	pSport1
HLXNE31	HLHDP51	HCRMY95	НАQBZ89	HYAAS90	HLDCP20	HWLJS42	HWLEH32	HWLGI62	HWLEL81
		001	70		92			49	97
		100	49		92			46	97
1288	279	116	430	341	1043	243	096	850	959
1043	55	3	2	Ξ	3		793	272	3
4516	4517	4518	4519	4520	4521	4522	4523	4524	4525
		gb AAC33514.	gb AAB37999. 1		emb CAB6647 8.1			gb AAC19403.	gb AAD48398. 1 AF1270
		(AF019767) zinc finger protein [Homo sapiens] >sp 075312 ZPR1_HUMAN ZINC-FINGER PROTEIN ZPR1. Length = 459	strong similarity to class-III of pyridoxal-phoshate-dependent aminotransferases [Caenorhabditis elegans] >pir T25848 T25848 hypothetical protein T01B11.2 - Caenorhabditis elegans >sp P91408 YO1J_CAEEL PROBABLE AMINOTRANSFERASE T01B11.2 (EC 2.6.1). Leng		(AL136543) hypothetical protein [Homo sapiens] >emb CAB66478.1  (AL136543) hypothetical protein [Homo sapiens] >sp CAB66478 CAB66478 Hypothetical 84.8 kd protein. >pir B34461 B34461 heat shock protein 90 beta - rabbit (fragment) {SUB 1-25} >sp P30947 HS9B			non-muscle myosin heavy chain [Bos taurus] >sp 002717 002717 NON-MUSCLE MYOSIN HEAVY CHAIN (FRAGMENT). Length = 625	(AF127035) calcium-activated chloride channel protein 2 [Homo sapiens] >sp[AAD48398]AAD48398 Calcium-activated chloride channel protein 2. >dbj BAA90969.1  (AK000138) unnamed protein product [Homo sapiens] {SUB 449-917} Length = 917
829084	829148	829161	830123	830151	830194	830231	830316	830343	830347
239	240	241	242	243	244	245	246	247	248

pCMVSport 3.0	pCMVSport 3.0	HUVDZ54 Uni-ZAP XR	pSport1	Uni-ZAP XR	pBluescript	Uni-ZAP XR	pCMVSport 2.0
HWHPA71	HWABR83	HUVDZ54	HUFAR83	HTLHR67	HTSGO78	HSLHS76	HKACP86
	86	94		100	84	87	94
	97	94		100	84	84	94
229	523	1319	219	733	1874	712	1192
2	83	51	420	2	E.	107	2
4526	4527	4528	4529	4530	4531	4532	4533
	dbj BAA74879 .1	emb CAA0150 3.1		gb AAB61533. 1	gb AAA18019. 1	emb CAB4330 9.1	dbj BAA00047 .1
	(AB020663) KIAA0856 protein [Homo sapiens] >sp O94938 O94938 KIAA0856 PROTEIN (FRAGMENT). Length = 1070	preA-PAI-2 [synthetic construct]   emb[CAA0150   >emb[CAA00247.1   miniactivin [synthetic construct]   3.1   {SUB 20-434} >gb AAA60005.1   plasminogen activator inhibitor 2, (first expressed exon) [Homo sapiens] {SUB 20-75} Length = 434		protein kinase MUK2 [Rattus norvegicus] >gb AAB95646.1  serine/threonine protein kinase [Rattus norvegicus] >sp P35465 PAK1_RAT SERINE/THREONINE-PROTEIN KINASE PAK-ALPHA (EC 2.7.1) (P68-PAK) (P21-ACTIVATED KINASE) (PAK-1) (ALPHA-PAK) (PROTEIN KINASE MU	tyrosine protein kinase [Homo sapiens] >sp Q08345 EDD1_HUMAN EPITHELIAL DISCOIDIN DOMAIN RECEPTOR 1 PRECURSOR (EC 2.7.1.112) (TYROSINE- PROTEIN KINASE CAK) (CELL ADHESION KINASE) (TYROSINE KINASE DDR) (DISCOIDIN RECEPTOR TYROSINE KINASE) (TRK E) (PROTEIN-T		(2"-5")oligoadenylate synthetase [Homo sapiens] Length = 364
830382	830436	830465	830498	830540	830568	830582	830586
249	250	251	252	253	254	255	256

pCMVSport 3.0	pSport1	Uni-ZAP XR	HODGW05 Uni-ZAP XR	pCMVSport 3.0	pSport1
HWBEJ14	HVAAB82	HPWBX45	HODGW05	HNTCW73	HASAB03
82	100	96	99	89	94
08	100	94	53	45	94
735	592	1005	235	1106	1697
	2	190	297	r	m
4542	4543	4544	4545	4546	4547
gb AAF04618. 1 AF0973	gb AAF21240. 1 AF0039	.1	gb AAA58464. 1	gb AAA28097. 1	emb CAA3079 0.1
(AF097362) gamma-interferon inducible lysosomal thiol reductase [Homo sapiens] >sp AAF04618 AAF04618 Gamma-interferon inducible lysosomal thiol reductase. >gb AAD22672.1 AC007192_3 (AC007192) INIP_HUMAN [AA 4-104] [Homo sapiens] {SUB 4-104} Length = 261	(AF003924) ANC_2H01 [Homo sapiens] >sp AAF21240 AAF21240 ANC_2H01. Length = 485	GLY1 protein [Escherichia coli] >dbj BAA20882.1  (AB005050) threonine aldolase [Escherichia coli] >gb AAC73957.1  (AE000188) putative arylsulfatase [Escherichia coli] >pir F64825 F64825 L-allothreonine aldolase (EC 4.1.2) - Escherichia coli >sp P75823	ORF 3 [Homo sapiens] >pir E41925 E41925 hypothetical protein 3 - human >sp Q14270 Q14270 ORF 3. Length = 143	coded for by C. elegans cDNAs GenBank: CE5D1 (Z14791), CEL01F1 (M88817), CEL04B5(M88849), and CEL04C1(M75812); putative [Caenorhabditis elegans] >pir S44853 S44853 K12H4.3 protein - Caenorhabditis elegans >sp P34524 YM63_CAEEL HYPOTHETICAL 40.2 KD PROTEIN	integrin beta 1 subunit precursor [Homo sapiens] >pir B27079 B27079 fibronectin receptor beta chain precursor - human >sp P05556 ITB1_HUMAN FIBRONECTIN RECEPTOR BETA_SUBUNIT PRECURSOR (INTEGRIN BETA-1) (CD29) (INTEGRIN VLA-4 BETA SUBUNIT). >gb AAA79835.1
830859	830879	830901	831019	831057	831099
265	266	267	268	269	270

271	831117			4548	400	579			HMWBR70	HMWBR70 Uni-ZAP XR
272	831163	IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593	sp P39194 AL U7_HUMAN	4549	3	161	75	83	HMSHS44	HMSHS44 Uni-ZAP XR
273	831210	TGF-beta masking protein large subunit [Rattus norvegicus] >pir[A38261 A38261 masking protein 1  precursor - rat >sp Q00918 TGFB_RAT LATENT TRANSFORMING GROWTH FACTOR BETA BINDING PROTEIN 1 PRECURSOR (TRANSFORMING GROWTH FACTOR BETA-1 BINDING PROTEIN 1) (TGF	gb AAA42235. 1	4550	-	498	98	91	HME1J62	Lambda ZAP II
274	831212	(AF051882) carbonic anhydrase XII precursor [Homo sapiens] >gb AAC63952.1  (AF037335) 1  carbonic anhydrase precursor [Homo sapiens] >sp O43570 CAHC_HUMAN CARBONIC ANHYDRASE XII PRECURSOR (EC 4.2.1.1) (CARBONATE DEHYDRATASE XII) (CA-XII) (TUMOR ANTIGEN HOM-R	gb AAC39789.	4551	206	385	100	100	НWННW79	pCMVSport 3.0
275	831234			4552	202	498			HLYGG06	pSport1
276	831239			4553	420	638			HMEKY46	HMEKY46 Lambda ZAP
277	831268			4554	324	749			HLTER57	HLTER57 Uni-ZAP XR
278	831307	(AK001317) unnamed protein product [Homo db sapiens] Length = 481	dbj BAA91619 .1	4555	33	935	94	94	HAPOA59	HAPOA59 Uni-ZAP XR
279	831313	555) [Mus musculus] transforming protein fos - FBJ virus VFB P55-V-FOS ROTEIN. Length = 381	emb CAA2450	4556	1182	1670	73	88	HAGDZ30	HAGDZ30 Uni-ZAP XR
280	831386			4557	1097	1363			HKLRB18	pBluescript

pSport1	pCMVSport 2.0	pSport1	ni-ZAP XR	ambda ZAP II	pCMVSport 3.0	Uni-ZAP XR
HKGDF04	HKAJZ24 po	HWLJE49	HJPAU37 Uni-ZAP XR	HHGCU20 Lambda ZAP	ннеро80 р	HFPCU40 U
94	94	83	93		8	72
94	93	83	06		94	72
1312	827	315	529	410	850	1574
254	3		2	3	2	180
4558	4559	4560	4561	4562	4563	4564
gb AAA51711. 1	emb CAB6510 4.1	gb AAA35886. 1	dbj BAA31675 .1		gb AAF14559. 1 AF1798	gb AAB20770. 1
aldehyde reductase (EC 1.1.1.2) [Homo sapiens] >gb AAB92369.1  (AF036683) aldehyde reductase [Homo sapiens] >gb AAF01260.1 AF112485_1 (AF112485) aldehyde reductase [Homo sapiens] >pir A33851 A33851 alcohol dehydrogenase (NADP+) (EC 1.1.1.2) - human >sp AA	(AJ245539) GalNAc-T5 [Homo sapiens] >sp CAB65104 CAB65104 GalNAc-T5 (fragment). Length = 668	2A9 peptide [Homo sapiens] >gb AAA51905.1  calcyclin [Homo sapiens] >gb AAA51906.1  put. calcyclin; putative [Homo sapiens] >pir A28363 BCHUY calcyclin - human >sp P06703 S106_HUMAN CALCYCLIN (PROLACTIN RECEPTOR ASSOCIATED PROTEIN) (PRA) (GROWTH FACTOR-IN	(AB014600) KIAA0700 protein [Homo sapiens] >sp 075182 075182 KIAA0700 PROTEIN (FRAGMENT). Length = 1130		(AF179867) STE20-like kinase [Homo sapiens] >sp AAF14559 AAF14559 STE20-like kinase. Length = 898	nuclear ribonucleoprotein complex K, man, Peptide, 463 aa] [Homo sapiens] 66.1   dC-stretch binding protein 1s norvegicus] >pir A42058 A42058 s nuclear ribonucleoprotein complex K, man
831390	831426	831453	831465	831558	831586	831664
281	282	283	284	285	286	287

	_	C- 4	1	1	1.22	T-4		· · · · ·				
HFKHD75 Uni-ZAP XR	pSport1	Uni-ZAP XR	HETBE76 Uni-ZAP XR	HTXOJ32 Uni-ZAP XR	Uni-ZAP XR	HE6FT69 Uni-ZAP XR	pCMVSport 2.0	pCMVSport 2.0	HLHGG05 Uni-ZAP XR	pCMVSport 3.0	pCMVSport 3.0	pSport1
	HFIHX78	HETIK68	HETBE76	HTXOJ32	HE9RY54	HE6FT69	HDTBQ51	HDTAB33	HLHGG05	HDPTH11	HDPLB15	HDAAQ89
100										77		
86										72		
305	240	219	1208	1855	1158	444	693	308	464	763	269	346
09	52	1	915	1490	892	-	-	3	201	44	3	2
4565	4566	4567	4568	4569	4570	4571	4572	4573	4574	4575	4576	4577
emb CAA5404 7.1										emb CAB4685 2.1		
hMpv17 [Homo sapiens] >glAAD14014.1 1683146_1 [Homo sapiens] >pir S45343 S45343 glomerulosclerosis protein Mpv17 - human >sp P39210 MPV1_HUMAN MPV17 PROTEIN. >gb AAC24205.1  (AF038633) Mpv17 protein [Homo sapiens] {SUB 155-176} Length = 176										(AJ388553) hypothetical protein [Canis familiaris] >sp[Q9XSR5 Q9XSR5 HYPOTHETICAL 15.3 KD PROTEIN (FRAGMENT). Length = 146		
831687	831703	831753	831757	831795	831796	831880	831899	831910	831931	831942	831956	832009
288	289	290	291	292	293	294	295	296	297	298	299	300

pCMVSport	pSport1	pSport1.	Uni-ZAP XR	pSport1	HCQAI40 Lambda ZAP	HWACZ95 pCMVSport	pSport1	pBluescript	pCMVSport 3.0	pBluescript SK-
HDFUB44	HGC0L40	HCRNJ73	HODEY51	HFIHN81	HCQAI40	HWACZ95	HBAGU45	HRGSB33	HAJBC35	H2LAJ21
69	66					77				66
57	66					09				66
348	1794	422	466	380	400	1487	471	489	360	1203
	-	279	317	246	26	462	295	196	145	49
4578	4579	4580	4581	4582	4583	4584	4585	4586	4587	4588
emb CAA1727 8.1	dbj BAA21762   51					emb CAB0159 0.1				gb AAC50062. 1
(AL021918) b34I8.1 (Kruppel related Zinc Finger protein 184) [Homo sapiens] >sp O60792 O60792 B34I8.1 (KRUPPEL RELATED ZINC FINGER PROTEIN 184). Length = 751	5-aminoimidazole-4-carboxamide ribonucleotide transformylase [Homo sapiens] >dbj BAA11559.1  5-aminoimidazole-4-carboxamide-1-beta-D-ribonucl eotide transformylase/inosinicase [Homo sapiens] >pir JC4642 JC4642 purH bifunctional enzyme - human >sp Q13856 Q					Similarity to E.coli 2-oxoglutarate dehydrogenase (SW:ODO1_ECOLI); cDNA EST EMBL:D32590 comes from this gene; cDNA EST EMBL:D32841 comes from this gene; cDNA EST EMBL:D34051 comes from this gene; cDNA EST EMBL:D35268 comes from this gene; cDNA >>pir[T2803				protein tyrosine kinase [Homo sapiens] >pir A55922 A55922 tyrosine kinase A6 - human >sp Q12792 Q12792 PROTEIN TYROSINE KINASE. Length = 350
832010	832044	832093	832138	832148	832187	832343	832346	832411	832464	832575
301	302	303	304	305	306	307	308	309	310	311

832593	93	CENP-F kinetochore protein [Homo sapiens] >sp P49454 CENF_HUMAN CENP-F KINETOCHORE PROTEIN. Length = 3210	gb AAA82889. 1	4589	2	811	91	92	H2LAB53	pBluescript SK-
832	832597			4590	214	318			H2CBJ07	pBluescript SK-
837	834890	TRANSCRIPTION FACTOR BTF3 (RNA pp[Q64152]B POLYMERASE B TRANSCRIPTION FACTOR 3). F3_MOUSE Length = 204	sp Q64152 BT F3_MOUSE	4591	70	588	87	88	H2CBT12	pBluescript SK-
83.	835079			4592	151	348			ноегн62	HOELH62 Uni-ZAP XR
83	835456	(AL035608) dJ479J7.1 (similar to CHONDROMODULIN-1) [Homo sapiens] >sp CAB55680 CAB55680 DJ479J7.1 (similar to CHONDROMODULIN-1) (fragment). Length = 263	emb CAB5568 0.1	4593	88	1041	79	79	HE8NG02	Uni-ZAP XR
83	835655			4594	1075	1332			HAGFG91	HAGFG91 Uni-ZAP XR
83	836203			4595	550	066			9/DOTMH	pSport1
83	836261	(AF117615) heme-binding protein [Homo sapiens] >sp Q9Y5Z5 Q9Y5Z5 HEME-BINDING PROTEIN. Length = 189	gb AAD32098. 1 AF1176	4596	116	292	86	86	HBMAD50	pBluescript SK-
83	836762	(AF132552) BcDNA.GM01838 [Drosophila melanogaster] >splQ9XZ53 Q9XZ53 BCDNA.GM01838. Length = 774	gb AAD27851. 1 AF1325	4597	2	1075	75	84	H2CBN10	pBluescript SK-
<b>∞</b>	836988	(AB011176) KIAA0604 protein [Homo sapiens] >sp O60344 ECE2_HUMAN ENDOTHELIN-CONVERTING ENZYME 2 (EC 3.4.24.71) (ECE-2) (KIAA0604). Length = 765	dbj BAA25530 .1	4598	68	571	87	87	HCE3J64	Uni-ZAP XR
83	838140			4599	300	476			НЕ2СН58	HE2CH58 Uni-ZAP XR
82	838459	!!!! ALU SUBFAMILY J WARNING ENTRY !!!! Length = 591	sp P39188 AL U1_HUMAN	4600	1223	1354	89	9/	HTHCW70	Uni-ZAP XR

Uni-ZAP XR	HTGEX11 Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	HTGAZ34 Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	Uni-ZAP XR
HAPOF13	HTGEX11	HWHGE39	HNGIN84	HTGAZ34	HNTEF54	HTEAF73	HPJCI42
66	93	100		95	99		93
66	92	100		95	99		93
1216	1069	1155	419	1172	592	1487	1044
26	5	-	09	m	2	1200	151
4601	4602	4603	4604	4605	4606	4607	4608
gb AAC39540.	emb CAB6374	gb AAD43021. 1		gb AAA19873. 1	emb CAB5591 9.1		emb CAB5527 8.1
(AF000364) heterogeneous nuclear ribonucleoprotein R [Homo sapiens] >pir T02673 T02673 heterogeneous nuclear ribonucleoprotein R - human >sp O43390 O43390 HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN R. Length = 633	(AL133623) hypothetical protein [Homo sapiens] >emb CAB63749.1  (AL133623) hypothetical protein [Homo sapiens] >sp CAB63749 CAB63749 Hypothetical 130.1 kd protein (fragment). Length = 1159	(AF100757) COP9 complex subunit 4 [Homo sapiens] >sp Q9Y677 Q9Y677 COP9 COMPLEX SUBUNIT 4. Length = 405		putative [Homo sapiens] >pir I54339 I54339 protoncogene bmi-1 - human >sp P35226 BMI1_HUMAN DNA-BINDING PROTEIN BMI-1. Length = 326	[Homo sapiens] hypothetical 229[T17229 156.1 - human thetical 39.8 kd		(AL035461) dJ967N21.6 (novel CDP-alcohol phosphatidyltransferase family member protein) [Homo sapiens] >sp CAB55278 CAB55278 DJ967N21.6 (novel CDP-alcohol phosphatidyltransferase family member protein). Length = 301
839262	839384	839750	840028	840572	840675	840708	840847
324	325	326	327	328	329	330	331

332	840848	sapiens] Length =	gb AAB21614.	4609	81	917	93	93	ннвнм75	HHBHM75 pCMVSport
333	840860	NAP [Homo sapiens] >pir S40510 S40510 nucleosome assembly protein 1-like 1 - human >sp P55209 NPL1 HUMAN NUCLEOSOME ASSEMBLY PROTEIN 1-LIKE 1 (NAP-1 RELATED PROTEIN). Length = 391	gb AAC37544. 1	4610	92	1309	89	89	HDTLJ39	pCMVSport 2.0
334	841015			4611	48	425			HE2DT31	Uni-ZAP XR
335	841017			4612	402	683			HE2AY01	Uni-ZAP XR
336	841030			4613	515	721			HWLOA34	pSport1
337	841241	Thy-1 [Homo sapiens] >pir A02106 TDHU Thy-1 membrane glycoprotein precursor - human >sp P04216 THY1_HUMAN THY-1 MEMBRANE GLYCOPROTEIN PRECURSOR (THY-1 ANTIGEN) (CDW90) (CD90 ANTIGEN). Length = 161	gb AAA61180. 1	4614	128	622	98	87	HBXFG67	HBXFG67 ZAP Express
338	841957			4615	355	609			HWLOF51	pSport1
339	846025	(AJ010973) DEDD protein [Homo sapiens] >gb AAC33105.1  (AF083236) FLDED-1 [Homo sapiens] >gb AAC80280.1  (AF043733) death effector domain-containing testicular molecule [Homo sapiens] >gb AAD16414.1  (AF100341) death effector domain-containing protein DED	emb CAA0944 5.1	4616	-	1098	63	83	HLDOK36	pCMVSport 3.0
340	846362	(AC006950) IgG Fc binding protein [AA 4671-5405] gb AAD15624. [Homo sapiens] >sp O95784 O95784 IGG FC   1  BINDING PROTEIN (FRAGMENT). Length = 735	gb AAD15624. 1	4617	449	1894	93	93	HSDJF12	Uni-ZAP XR
341	846384	(AF127036) calcium-activated chloride channel protein 1 [Homo sapiens] >sp[AAD25487]AAD25487 Calcium-activated chloride channel protein 1. Length = 914	gb AAD25487. 1 AF1270	4618	1125	2780	96	96	HWLFF02	pSport1

342	846750	(AF132148) [Drosophila melanogaster]   sp Q9XYZ6 Q9XYZ6 HYPOTHETICAL 75.5 KD   PROTEIN, Length = 653	gb AAD34736. 1	4619	1	1503	47	61	HEMFI21	HEMF121 Uni-ZAP XR
343	847289			4620	322	510			99MNTMH	pSport1
344	847598	IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593	sp P39194 AL U7_HUMAN	4621	710	859	85	87	HNTEG90	pCMVSport 3.0
345	848119	3cDNA.GH06193 [Drosophila   >sp Q9Y138 Q9Y138 06193. Length = 696	gb AAD38609. 1 AF1456	4622	_	243	99	88	HELGG49	HELGG49 Uni-ZAP XR
346	848746			4623	250	999			HWLQ044	pSport1
347	849084	ATP synthase subunit e [Homo sapiens] db >sp P56385 ATPJ HUMAN ATP SYNTHASE E .1  CHAIN, MITOCHONDRIAL (EC 3.6.1.34). {SUB 2-69} Length = 69	dbj BAA23322	4624	-	270	78	78	HFEBT64	Uni-ZAP XR
348	849114	epidermal growth factor receptor kinase substrate [Homo sapiens] >pir I38728 I38728 epidermal   I  growth factor receptor kinase substrate - human >sp Q12929 EPS8_HUMAN EPIDERMAL GROWTH FACTOR RECEPTOR KINASE SUBSTRATE EPS8. Length = 822	gb AAA62280. 1	4625	3	986	100	100	HUVFL24	HUVFL24 Uni-ZAP XR
349	849143	zinc finger protein PZF [Mus musculus] gb >pir[148724][48724 zinc finger protein PZF - mouse 1  >sp Q62511 Q62511 ZINC FINGER PROTEIN PZF. Length = 455	gb AAA81913. 1	4626	62	1795	88	68	HAMGR89	pCMVSport 3.0
350	849155	carbonic anhydrase I (EC 4.2.1.1) [Homo sapiens] gb >emb CAA28663.1  carbonic anhydrase I (AA 1-261) [Homo sapiens] >pir JQ0786 CRHU1 carbonate dehydratase (EC 4.2.1.1) I - human >sp P00915 CAH1_HUMAN CARBONIC ANHYDRASE I (EC 4.2.1.1) (CARBONATE DEHYDRATASE	gb AAA51910. 1	4627	71	859	100	100	HKLSA58	pBluescript

pSport1	Uni-ZAP XR	pCMVSport 3.0	pBluescript SK-	pBluescript	Uni-ZAP XR	Uni-ZAP XR	Uni-ZAP XR
HWLCG11	HMSJT69	HRABQ68	H2CBM53	HPRTG34	HE8DO31	HAIDB85	HMCIR67
66	100		100	68	66	100	95
66	100		100	68	66	100	95
903	877	323	1561	534	831	933	1788
<b></b>	110	18	164	-	13	-	889
4628	4629	4630	4631	4632	4633	4634	4635
emb CAA6695 5.1	dbj BAA91753		gb AAA59967. 1	dbj BAA91207 .1	dbj BAA34595 .1	emb CAB5307	dbj BAA07011 .1
carcinoembryonic antigen [Homo sapiens] emb CAA6695   >gb AAA66186.1  carcinoembryonic antigen [Homo sapiens] >gb AAC62825.1  (AC005797)   carcinoembryonic antigen CGM2 precursor - human [Homo sapiens] >pir A55811 A55811   carcinoembryonic antigen CGM2 precursor - human >s	(AK001553) unnamed protein product [Homo sapiens] >dbj BAA91996.1  (AK001951) unnamed protein product [Homo sapiens] Length = 227		ornithine decarboxylase [Homo sapiens] >gb AAA59969.1  ornithine decarboxylase [Homo sapiens] >gb AAA60563.1  ornithine decarboxylase [Homo sapiens] >gb AAA60564.1  ornithine decarboxylase [Homo sapiens] >emb CAA39047.1  ornithine decarboxylase [Homo sapi		5T:RICS2753A [Homo 95571 MRNA EXPRESSED 7. Length = 227	(AL035071) dJ1085F17.2 (EB1 (APC binding protein)) [Homo sapiens] >gb AAC09471.1  EB1 [Homo sapiens] >pir I52726 I52726 EB1 - human >sp Q15691 Q15691 EB1.  >sp CAB53072 CAB53072 DJ1085F17.2 (EB1 (APC binding protein)). Length = 268	thromboxane synthase [Homo sapiens] Length = 533   dbj BAA07011   .1
849159	849244	849254	849301	849317	849332	849422	849471
351	352	353	354	355	356	357	358

pCMVSport 2.0	pSport1	HPRAO21 Uni-ZAP XR	Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	Uni-ZAP XR
HKAJC79	HCRMP14	HPRAO21	HAIBU93	HCFMH52	HMVAE41	HMSDT39 Uni-ZAP XR	HE8NK24
100	100		9/	100	53	66	95
100	100		89	66	47	86	95
595	208	1036	1851	508	896	1100	1169
2	302	908	-	44	831	57	8
4636	4637	4638	4639	4640	4641	4642	4643
gb AAA31492. 1	gb AAA67217.		gb AAF18302. 1 AF1202	dbj BAA23735 .1	gb AAB97675.	gb AAA59570. 1	emb CAA0583 2.1
ubiquitin conjugating-protein [Oryctolagus cuniculus] >gb AAA35982.1  HHR6B (Human homologue of yeast RAD 6); putative [Homo sapiens] >emb CAA37339.1  E2 protein [Homo sapiens] >gb AAA21087.1  ubiquitin conjugating-protein [Rattus norvegicus] >emb CAA6560	beta-galactosidase alpha peptide [Cloning vector pSport2] Length = 114		(AF120206) XY body protein [Mus musculus]   >gb AAF18303.1 AF120207_1 (AF120207) XY body protein [Mus musculus]   >sp AAF18302 AAF18302 XY body protein.   >sp AAF18303 AAF18303 XY body protein.   = 840	(AB009282) cytochrome b5 [Homo sapiens] >sp O43169 O43169 CYTOCHROME B5 (FRAGMENT). Length = 146	(AF020762) protein [Homo sapiens] >sp O43466 O43466 HYPOTHETICAL 31.3 KD PROTEIN (FRAGMENT). Length = 267	macrophage capping protein [Homo sapiens] >pir A43358 A43358 macrophage capping protein - human >sp P40121 CAPG_HUMAN MACROPHAGE CAPPING PROTEIN (ACTIN- REGULATORY PROTEIN CAP-G). >gb AAA92670.1  Cap-G [Homo sapiens] {SUB 1- 172} Length = 348	(AJ003061) most expressed alternative spliced form [Homo sapiens] >splO60852 O60852 PROTEIN ENCODED BY SACCHAROMYCES CEREVISIAE SPC98 HOMOLOGUE. Length = 907
849492	849534	849565	849583	849589	849658	849666	849679
359	360	361	362	363	364	365	366

pCMVSport 3.0	pSport1	HTOAC26 Uni-ZAP XR	HUVCQ41 Uni-ZAP XR	Uni-ZAP XR	HCQCD86 Lambda ZAP	pSport1	Uni-ZAP XR	pCMVSport 3.0	pSport1	Uni-ZAP XR	pSport1	pCMVSport 2.0
нwноро8	HCRPJ23	HTOAC26	HUVCQ41	нрлесе6	Э8СОСО86	HCRMX05	HAPRB43	нwнqг22	HWLMN93	HTGFW53	HANGG89	HKAAV86
	95		87	26			100	86		74		100
	95		80	26			66	86		73		100
1067	1127	213	1144	2145	476	182	513	1033	273	1681	496	852
948	69	52	1031	601	336	69	1	434	1	1550	569	1
4644	4645	4646	4647	4648	4649	4650	4651	4652	4653	4654	4655	4656
	gb AAA91639. 1		sp P39194 AL U7_HUMAN	dbj BAA19764 .1			gb AAD17317.	dbj BAA91068 .1		sp P39194 AL U7_HUMAN		emb CAA7373 4.1
	lumican [Homo sapiens] Length = 338		IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593	a-D-glucosaminide [Homo sapiens] 1985.1 glycoprotein 6-alpha-L- srase [Homo sapiens] 1986.1 glycoprotein 6-alpha-L- srase [Homo sapiens] 1987.1 glycoprotein 6-alpha-L- srase [Ho			(AF124522) tetraspan NET-2 [Homo sapiens] gb AAD17317. >sp O95859 O95859 TETRASPAN NET-2. Length = 1  305	(AK000302) unnamed protein product [Homo sapiens] Length = 436		IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593		GDP dissociation inhibitor beta [Homo sapiens] >emb CAA73735.1  GDP dissociation inhibitor beta [Homo sapiens] >sp O43928 O43928 GDP DISSOCIATION INHIBITOR BETA. >gb AAD34588.1  (AF144713) Rab GDP dissociation inhibitor beta [Homo sapiens] {SUB 81-439} Le
849741	849783	850211	850254	850264	850273	850371	850859	851066	851217	852170	852387	852812
367	368	369	370	371	372	373	374	375	376	377	378	379

380	853175	III! ALU SUBFAMILY SQ WARNING ENTRY   III! sp P39194 AL   Length = 593	sp P39194 AL U7_HUMAN	4657	234	809	82	84	HSACF33	pBluescript SK-
381	853230			4658	19	273			H2CBA56	pBluescript SK-
382	854063			4659	-	129		_	HLJBL63	pCMVSport 1
383	854073	(AF068754) heat shock factor binding protein 1 HSBP1 [Homo sapiens] >sp O75506 O75506 HEAT SHOCK FACTOR BINDING PROTEIN 1 HSBP1. Length = 76	gb AAC25186. 1	4660	146	403	100	100	HHFOV83	Uni-ZAP XR
384	854987	no arches [Homo sapiens] >sp 095639 095639 NO ARCHES. Length = 269	gb AAD00321. 1	4661	9/	408	100	100	HMTAE04	pCMVSport 3.0
385	855130			4662	127	333			HWLNN76	pSport1
386	856227	arrestin [Homo sapiens] >pir S18984 S18984 arrestin emb CAA7757 - human (fragment) >sp P32121 ARR2_HUMAN 7.1  BETA-ARRESTIN 2. Length = 409	emb CAA7757	4663	-	636	59	62	нроғе56	pCMVSport 3.0
387	856243	similar to citrate Jyase beta chain; cDNA EST yk302b4.5 comes from this gene [Caenorhabditis elegans] >pir[718818 T18818] hypothetical protein C01G10.7 - Caenorhabditis elegans >sp Q93167 Q93167 C01G10.7 PROTEIN. Length = 324	emb CAB0270	4664	-	411	35	54	HLDBR21	pCMVSport 3.0
388	856354	KIAA0176 [Homo sapiens] >sp Q14681 Y176_HUMAN HYPOTHETICAL PROTEIN KIAA0176 (FRAGMENT). Length = 265	dbj BAA11493 .1	4665	-	591	40	95	ННАОДЭ1	pCMVSport 3.0
389	856923	(AF132951) CGI-17 protein [Homo sapiens] >sp Q9Y306 Q9Y306 CGI-17 PROTEIN. Length = 385	gb AAD27726. 1 AF1329	4666	<del>ر</del>	881	86	86	<b>НТОНА37</b>	HTOHA37 Uni-ZAP XR
390	857684			4667	166	2			HDPPP71	pCMVSport 3.0

391	857946	(AF077034) HSPC010 [Homo sapiens] >gb AAD33954.1 AF145385_1 (AF145385) hypoxia-inducbile gene 1 [Homo sapiens] >sp Q9Y241 Q9Y241 HIG1 PROTEIN (HSPC010). Length = 93	gb AAD27767. 1 AF0770	4668	_	408	100	100	HBBBE52	HBBBE52 pCMVSport
392	858166			4669	641	1006			HLTDR01	HLTDR01 Uni-ZAP XR
393	858178	KIAA0183 [Homo sapiens] >sp Q14688 Q14688 KIAA0183 PROTEIN (FRAGMENT). Length = 1062	dbj BAA11500	4670	1837	3153	87	87	НМЕСD50	HMECD50 Lambda ZAP
394	858606			4671	2	307			HDPJL40	pCMVSport 3.0
395	858894	KIAA0021 protein [Homo sapiens] >sp BAA03499 BAA03499 KIAA0021 protein (fragment). Length = 703	dbj BAA03499 .2	4672	2	1894	68	68	HDPGS38	pCMVSport 3.0
396	858949			4673	29	322			нсоам69	HCQAM69 Lambda ZAP
397	828928			4674	2	106			HOSNC15	HOSNC15 Uni-ZAP XR
398	859171	Similar to Human C219-reactive peptide (L34688) [Homo sapiens] >sp Q92580 Q92580 MYELOBLAST KIA0268 (FRAGMENT). >sp AAB00324.1 C219-reactive peptide [Homo sapiens] {SUB 592-727} Length = 1193	dbj BAA13448 .1	4675	8	1463	84	84	ннело41	pCMVSport 3.0
399	859352	(AF107406) GW128 [Homo sapiens] >sp Q9Y649 Q9Y649 GW128. Length = 63	gb AAD44524. 1 AF1074	4676	177	398	100	100	HTXMR51	HTXMR51 Uni-ZAP XR
400	859354	(AK001214) unnamed protein product [Homo sapiens] Length = 532	dbj BAA91559 .1	4677	3	1244	66	66	ннғсх08	HHFCX08 Uni-ZAP XR
401	859702			4678	99	310			HNTEG54	pCMVSport 3.0

	92 HCDEA29 Uni-ZAP XR	HCYBJ35 pBluescript SK-	96 HEBGA63 Uni-ZAP XR	HFACI10 Uni-ZAP XR	96 HETCM67 Uni-ZAP XR	HCRNF78 pSport1	HRACX96 pCMVSport 3.0	90 HTLAK94 Uni-ZAP XR	HCQCV31 Lambda ZAP
100	91		96	,	96			06	
929	1773	2	1029	841	1318	194	144	1706	417
321	91	115	103	311	2	15	1	654	256
4679	4680	4681	4682	4683	4684	4685	4686	4687	4688
gb AAC39530. 1	gb AAC51244. 1		gb AAA36352. 1		emb CAB4208 5.1			gb AAD34047. 1 AF1518	
(AF010312) Pig7 [Homo sapiens] >gb AAB36550.1  gb AAC39530. LPS-Induced TNF-Alpha Factor [Homo sapiens] 1  >sp Q99732 LITF_HUMAN LIPOPOLYSACCHARIDE-INDUCED TUMOR NECROSIS FACTOR-ALPHA FACTOR (LPS-INDUCED TNF-ALPHA FACTOR) (P53 INDUCED PROTEIN 7). Length = 228	collagen type XII alpha-1 [Homo sapiens] >sp Q99715 CA1C_HUMAN COLLAGEN ALPHA 1(XII) CHAIN PRECURSOR. >pir A44036 A44036 collagen alpha 1(XII) chain - bovine (fragment) {SUB 2492-2517} Length = 3063		Na,K-ATPase beta subunit [Homo sapiens] >emb CAA27385.1  put. Na/K-ATPase beta (aa 1-303) [Homo sapiens] >pir A23764 PWHUNB Na+/K+-exchanging ATPase (EC 3.6.1.37) beta chain - human >sp P05026 ATNB_HUMAN SODIUM/POTASSIUM-TRANSPORTING ATPASE BETA-1 CHAIN (		(AJ242015) eMDC II protein [Homo sapiens] >sp Q9Y3S0 Q9Y3S0 EMDC II PROTEIN. Length = 775			(AF151810) CGI-52 protein [Homo sapiens] >sp Q9Y365 Q9Y365 CGI-52 PROTEIN. Length = 359	
860915	861209	861534	861697	861826	861909	862197	862232	862237	862277
402	403	404	405	406	407	408	409	410	411

HTJMG70 pCMVSport 2.0	52 Uni-ZAP XR	HHFCZ67 Uni-ZAP XR	HHFIA95 Uni-ZAP XR	85 Uni-ZAP XR	8 Uni-ZAP XR	99 pCMVSport 3.0	49 pSport1	46 pCMVSport 3.0	67 pCMVSport 3.0	45 pBluescript	PCMVSport 3.0	81 pSport1	52 Uni-ZAP XR
HTJMG	HSNAT52	HHFCZ	HHFIA9	HMSOR85	HBJJU68	HDPBN09	HFNAC49	HHETS46	HHATS67	HLHTL45	HHEJZ45	HNAAF81	HSLGX52
			68			19				_		6	
			88			46						6	
727	185	880	1178	493	207	1385	972	574	447	485	969	2247	613
533	93	200	192	323	-	К	151	209	256	171		253	359
4689	4690	4691	4692	4693	4694	4695	4696	4697	4698	4699	4700	4701	4702
			dbj BAA19546 .1			emb CAA9436 8.1						dbj BAA91687  .1	
			(AB002533) Qip1 [Homo sapiens] >gb AAC25605.1  dbj BAA19546 importin alpha 3 [Homo sapiens] 11  2pirJUC5505 JUC5505 DNA helicase Q1 interacting protein 1 - human >sp O00629 IMA4_HUMAN IMPORTIN ALPHA-4 SUBUNIT (KARYOPHERIN ALPHA-4 SUBUNIT) (QIP1 PROTEIN). Length = 521			similar to Glutathione S-transferases.; cDNA EST yk536e7.3 comes from this gene [Caenorhabditis elegans] >pir[T24175 T24175 hypothetical protein R11A8.5 - Caenorhabditis elegans >sp Q21925 Q21925 R11A8.5 PROTEIN. Length = 347						(AK001431) unnamed protein product [Homo sapiens] Length = 597	
862285	862423	862456	862486	862709	863865	863944	864428	864808	864822	865044	865420	865421	866287
412	413	414	415	416	417	418	419	420	421	422	423	424	425

	+	~	~	<del> </del>	<u>~</u>	T	
pSport1	pCMVSport 2.0	H6EAB24 Uni-ZAP XR	Uni-ZAP XR	pCMVSport	HTAHC93 Uni-ZAP XR	Other	pSport1
HWLNL21	HKADX79	H6EAB24	HRDFP67	HDPPM58	нтансоз	HPCRL51	HCRNJ44
	96			100	40	93	74
	95			100	37	91	70
296	647	415	191	1749	461	703	851
671	3	80	3	16	8	473	336
4703	4704	4705	4706	4707	4708	4709	4710
	gb AAF34791. 1 AF2286			dbj BAA34600 .1	gb AAA21145. 1	emb CAB4330 6.1	gb AAD45423. 1 AF0991
	(AF228603) pleckstrin 2 [Homo sapiens] Length = 353			Asparaginyl tRNA Synthetase [Homo sapiens] >emb CAA04008.1  (AJ000334) asparaginyl-tRNA synthetase [Homo sapiens] >sp O43776 SYN HUMAN ASPARAGINYL- TRNA SYNTHETASE, CYTOPLASMIC (EC 6.1.1.22) (ASPARAGINE TRNA LIGASE) (ASNRS). >sp BAA34600 BAA34600 Asparag	AF-17 [Homo sapiens] >pir 138533 138533 AF17 protein - human >sp P55198 AF17 HUMAN AF-17 PROTEIN. >emb CAB63772.1  (AL133659) hypothetical protein [Homo sapiens] {SUB 670- 1093} Length = 1093	(AL050170) hypothetical protein [Homo sapiens] >emb CAB43306.1  (AL050170) hypothetical protein [Homo sapiens] >pir T08792 T08792 hypothetical protein DKFZp586E1422.1 - human (fragment) >sp Q9Y3Y1 Q9Y3Y1 HYPOTHETICAL 14.3 KD PROTEIN (FRAGMENT). Length = 1	(AF099186) EH domain-containing protein EHD1 [Mus musculus] >gb AAF24223.1 AF173156_1 (AF173156) MPAST1 [Mus musculus] >sp Q9WVK4 Q9WVK4 EH DOMAIN-CONTAINING PROTEIN EHD1.
866300	866414	286998	867132	867388	867842	867923	868035
426	427	428	429	430	431	432	433

200	BST-2 [Homo sapiens] >pir A56836 A56836 bone marrow stromal cell surface protein BST-2 - human >sp Q10589 BST2_HUMAN BONE MARROW STROMAL ANTIGEN 2 (BST-2). Length = 180	dbj BAA05679  .1	4711	78	959	82	82	HFKMJ43	Uni-ZAP XR
			4712	963	1169	-		HMSFS13	Uni-ZAP XR
			4713	1	615			нскон59	pSport1
(AK001655) unnamed prapiens] Length = 372	protein product [Homo	dbj BAA91815  -1	4714	2	1225	66	66	HHFJU87	Uni-ZAP XR
nuclear protein SA-2 [H >sp 000540 000540 NI Length = 1162	nuclear protein SA-2 [Homo sapiens] >sp 000540 000540 NUCLEAR PROTEIN SA-2. Length = 1162	emb CAA9973 2.1	4715	009	1541	93	93	HFIAUS9	pSport1
Ca2+ ATPase of fast-twitch skeletal mus sacroplasmic reticulum, adult isoform [H sapiens] >sp O14983 O14983 CA2+ ATF FAST-TWITCH SKELETAL MUSCLE SACROPLASMIC RETICULUM, ADU ISOFORM. Length = 1001	Ca2+ ATPase of fast-twitch skeletal muscle sacroplasmic reticulum, adult isoform [Homo sapiens] >sp O14983 O14983 CA2+ ATPASE OF FAST-TWITCH SKELETAL MUSCLE SACROPLASMIC RETICULUM, ADULT ISOFORM. Length = 1001	gb AAB53113. 1	4716	592	1452	88	68	HBKDR59	pSport1
!!!! ALU SUBFAMILY . Length = 591	Y J WARNING ENTRY !!!!	sp P39188 AL U1_HUMAN	4717	396	536	89	79	HTHCZ54	Uni-ZAP XR
ezrin (AA 1-586) [Homo sapiens]  >pir A34400 A34400 ezrin - human  >sp P15311 EZR1 HUMAN EZRIN (CYTOVILLIN) (VILLIN-2). {SUI  >gb AAA61278.1  cytovillin 2 [Hon  12-586} Length = 586	ezrin (AA 1-586) [Homo sapiens]  >pir A34400 A34400 ezrin - human  >sp P15311 EZRI_HUMAN EZRIN (P81)  (CYTOVILLIN) (VILLIN-2). {SUB 2-586}  >gb AAA61278.1  cytovillin 2 [Homo sapiens] {SUB 12-586} Length = 586		4718	79	1386	98	98	HWABV82	pCMVSport 3.0
(AF044286) histone ma >sp 075377 075377 HI Length = 369	(AF044286) histone macroH2A1.1 [Homo sapiens] >sp 075377 075377 HISTONE MACROH2A1.1. Length = 369	gb AAC33434. 1	4719	142	099	100	100	HACAC44	HACAC44 Uni-ZAP XR
			4720	378	629			HDTLE81	pCMVSport 2.0
			4721	12	158			HSWBU77	pCMVSport 3.0

CMVSport 3.0	pBluescript	pSport1	ni-ZAP XR	pSport1	pCMVSport 2.0	ni-ZAP XR	pBluescript	pSport1	pCMVSport 3.0	pCMVSport 3.0
HWACJ61 pCMVSport	HKLSC04 p	HCRPM84	HLHGG41   Uni-ZAP XR	HWLNH36	HKAAC09 p	HLHAR50   Uni-ZAP XR	HSKJB43 F	HNSMB24	HAJAN23 p	HWBAP55 p
94		66			88	77	100	57	08	001
94		66			88	29	86	42	64	100
802	2964	2	719	909	841	952	403	1108	1807	1954
2	2997	490	447	291	68	818	2	530	530	2
4722	4723	4724	4725	4726	4727	4728	4729	4730	4731	4732
gb AAC51860.		dbj BAA91694 .1			emb CAB7565 6.1	gb AAA40456. 1	dbj BAA23885 .1	gb AAC51784.	emb CAA7961 8.1	dbj BAA25472 .1
(AF029684) IkB kinase-beta [Homo sapiens] >gb AAD08997.1  (AF080158) IkB kinase-b [Homo sapiens] >sp O14920 O14920 IKB KINASE-BETA (FRAGMENT). Length = 756		(AK001443) unnamed protein product [Homo sapiens] Length = 420			(AL157432) hypothetical protein [Homo sapiens] Length = 221	11	RNA splicing-related protein [Rattus norvegicus] >sp 054729 054729 BRAIN. Length = 712	serine protease [Homo sapiens] >sp[O15393 TMS2_HUMAN TRANSMEMBRANE PROTEASE, SERINE 2 (EC 3.4.21). Length = 492	predicted using Genefinder; similar to Propionyl-CoA carboxylase beta chain; cDNA EST EMBL:M89018 comes from this gene; cDNA EST EMBL:D28069 comes from this gene; cDNA EST EMBL:D28068 comes from this gene; cDNA EST EMBL:D33966 comes from this >>pir S2831	(AB011118) KIAA0546 protein [Homo sapiens] >pir T00325 T00325 hypothetical protein KIAA0546 - human (fragment) >sp O60293 O60293 KIAA0546 PROTEIN (FRAGMENT). Length = 632
871225	871428	871498	871732	871756	871821	872327	872354	872535	872551	872640
445	446	447	448	449	450	451	452	453	454	455

HE2JO26 Uni-ZAP XR	HEGAK44 Uni-ZAP XR	pCMVSport 2.0	Uni-ZAP XR	pSport1	Uni-ZAP XR	Uni-ZAP XR	pSport1	pCMVSport 2.0	pSport1	pSport1	pCMVSport 3.0
HE2J026	HEGAK44	HOGCK09	НЕ9ҒН03	HWLUI05	HCEVS38	HE2BS79	HHMMB54	HKABZ52	HCR0J11	HWLJP34	HSYDL64
	95			100	71				98	26	81
	94			100	54				74	56	65
361	923	2150	505	513	564	384	293	689	37	767	503
2	192	1830	272	_	88	175	12	306	258	321	3
4733	4734	4735	4736	4737	4738	4739	4740	4741	4742	4743	4744
	dbj BAA91797 .1			emb CAB6479 2.1	gb AAC34490. 1				pir JU0033 JU 0033	gb AAD48816. 1 AF1641	dbj BAA20802 .1
	(AK001631) unnamed protein product [Homo sapiens] Length = 390			(AJ132860) receptor for activated C kinase [Bos taurus] >emb CAA83944.1  G-beta like protein [Sus scrofa] >gb AAD37978.1  (AF146043) RACK1 [Sus scrofa] >gb AAA59626.1  MHC B complex protein 12.3 [Homo sapiens] >emb CAA53062.1  B complex protein mRNA 12-3	(AC005313) protein [Arabidopsis thaliana] >pir[T02714 T02714 hypothetical protein T18E12.21 - Arabidopsis thaliana >sp O81062 O81062 T18E12.21 PROTEIN. Length = 344					(AF164119) CRIB-containing BORG3 protein [Mus musculus] >sp AAD48816 AAD48816 CRIB-containing BORG3 protein. Length = 150	(AB002342) KIAA0344 [Homo sapiens] >sp O15052 O15052 KIAA0344. Length = 1246
872655	872802	872852	873299	873633	874164	874307	874308	874309	874310	874320	874325
456	457	458	459	460	461	462	463	494	465	466	467

HCE1G78 Uni-ZAP XR	HSOBR31 Uni-ZAP XR	HLLCC54 pCMVSport	HE2LO76 Uni-ZAP XR	3 Uni-ZAP XR	7 pSport1	8 pSport1	HWADK27 pCMVSport	1 pSport1	4 pSport1
HCE1G78	HSOBR3	HLLCC5	HE2LO76	HTTIU53	HUFDS37	HWMAJ78	HWADK2	HCRNT71	HCRQA24
71	82					87		19	96
70	63					87		52	96
838	496	752	685	430	356	517	428	579	584
2	173	501	491	350	3	2	72	1	3
4745	4746	4747	4748	4749	4750	4751	4752	4753	4754
gb AAD15618. 1	gb AAD05037. 1					gb AAB97620. 1		gb AAF11464. 1 AE0020	dbj BAA86513 .1
(AC005005) similar to phosphatidylinositol (4,5)bisphosphate 5-phosphatase; match to PID:g1399105 [Homo sapiens] >sp AAD15618 AAD15618 WUGSC:H_DJ412A9.2 protein (fragment). Length = 1056	AMSH [Homo sapiens] >sp O95630 O95630 AMSH. gb AAD05037. Length = 424					(AC004030) F21856_2 [Homo sapiens] >pir T00636 T00636 hypothetical protein F21856_2 - human >sp O43360 O43360 F21856_2. Length = 679		(AE002030) thermoresistant gluconokinase [Deinococcus radiodurans] >pir B75338 B75338 thermoresistant gluconokinase - Deinococcus radiodurans (strain R1) >sp AAF11464 AAF11464 Thermoresistant gluconokinase. Length = 172	(AB033025) KIAA1199 protein [Homo sapiens] >sp BAA86513 BAA86513 KIAA1199 protein (fragment). Length = 1013
874327	874328	874329	874330	874345	874348	874349	874350	874352	874358
468	469	470	471	472	473	474	475	476	477

Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	pCMVSport 3.0	pSport1	pBluescript	Uni-ZAP XR	pSport1	pCMVSport 3.0	pSport1	Uni-ZAP XR
HUVCM45 Uni-ZAP XR	HRAAG89	HSLJR04	HNTBD52	HNTST27	HSKJH49	HOEMK72	HBKDS37	HJMAK37	HUSGS50	HTOIL45
98		88	08	88		·				77
98		84	74	82						77
1501	230	1449	335	472	210	1484	405	501	986	863
2	3	1787	<del>د</del>	2	31	105	190	76	619	က
4755	4756	4757	4758	4759	4760	4761	4762	4763	4764	4765
gb AAA82930.		dbjlBAA35615	gb AAF35182. 1 AF1957	gb AAB05478. 1						emb CAA0616
methionine aminopeptidase [Homo sapiens] >gb AAC63402.1  eIF-2-associated p67 homolog [Homo sapiens] >pir S52112 DPHUM2 methionyl aminopeptidase (EC 3.4.11.18) 2 - human >sp P50579 AMP2_HUMAN METHIONINE AMINOPEPTIDASE 2 (EC 3.4.11.18) (METAP 2) (PEPTIDASE		Cell division protein FtsK. [Escherichia coli] >dbj BAA35622.1  Cell division protein FtsK. [Escherichia coli] >gb AAC73976.1  (AE000191) cell division protein [Escherichia coli] >pir A64828 A64828 cell division protein ftsK - Escherichia coli >sp P46889	(AF195765) L2DTL protein [Homo sapiens] Length = 730	suppressor of hairless protein 1 [Xenopus laevis] >sp Q91880 Q91880 SUPPRESSOR OF HAIRLESS PROTEIN 1. Length = 501						(AJ004856) connexin31 [Homo sapiens] >gb AAC95471.1  (AF099730) connexin 31 [Homo sapiens] >gb AAD11816.1  (AF052692) connexin 31; gap junctional protein cx31 [Homo sapiens] >pir JE0274 JE0274 connexin 31 - human >sp O75712 CXB3_HUMAN GAP JUNCTION BETA-3
874362	874368	874369	874370	874372	874396	874399	874400	874401	874403	874407
478	479	480	481	482	483	484	485	486	487	488

HLTGR10 Uni-ZAP XR	pSport1	HCQBD69 Lambda ZAP	HATBE07 Uni-ZAP XR	HCQDD86 Lambda ZAP	pSport1	pSport1	pSport1	pSport1	HCQDC45 Lambda ZAP	pBluescript SK-	pSport1	pBluescript SK-	HTODN93 Uni-ZAP XR
HLTGR10	HWLQF84	нсоврее	HATBE07	нсорря6	HUCNE27	HCRNL83	HCRNJ94	HCROK63	нсорс45	HCYBG26	HCRNV56	HCYBL48	HTODN93
				76						96			
				88						96			
444	530	427	545	110	349	347	202	476	<i>L</i> 66	385	497	828	258
175	168	248	312	3	68	168	17	216	788	2	204	532	16
4766	4767	4768	4769	4770	4771	4772	4773	4774	4775	4776	4777	4778	4779
				86Y5 Y6Y5						gb AAD45243. 1 AF1186			
				IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414						(AF118637) feline leukemia virus subgroup C receptor FLVCR [Homo sapiens] >sp Q9Y5Y0 Q9Y5Y0 C-RECEPTOR. >dbj BAA91679.1  (AK001419) unnamed protein product [Homo sapiens] {SUB 277-555} Length = 555			
874410	874411	874413	874414	874416	874417	874422	874423	874424	874426	874427	874428	874432	874433
489	490	491	492	493	464	495	496	497	498	499	200	501	502

503	874435	(AL110261) hypothetical protein [Homo sapiens] >emb CAB53702.1  (AL110261) hypothetical protein [Homo sapiens] >pir/T14782[T14782] hypothetical protein DKFZp586B0621.1 - human (fragment) >sp CAB53702 CAB53702 Hypothetical 22.8 kd protein (fragment). Length	emb CAB5370 2.1	4780	316	009	86	86	HWLQK42	pSport1
504	874436			4781	45	197			HODDD101	Uni-ZAP XR
505	874437	anthracycline-associated resistance ARX [Homo gb AAC99992. sapiens] >gb AAD24434.1 AF110957_1 (AF110957) 1 SUMO-1 activating enzyme subunit 2 [Homo sapiens] >sp O95605 O95605 ANTHRACYCLINE-ASSOCIATED RESISTANCE ARX. Length = 640	gb AAC99992. 1	4782	E.	1367	91	91	HNTDB90	pCMVSport 3.0
909	874438			4783	872	1105			HFPBQ02	Uni-ZAP XR
507	874447			4784	236	541			HTXSK90	Uni-ZAP XR
508	874449			4785	866	1258			HTECD58	Uni-ZAP XR
509	874452	beta-galactosidase alpha peptide [Cloning vector pSport2] Length = 114	gb AAA67217.	4786	432	623	82	82	нwгон59	pSport1
510	874455			4787	6	176			ннерр22	pCMVSport 3.0
511	874458	(AF075724) [Legionella pneumophila] >sp 085769 085769 HYPOTHETICAL 24.8 KD PROTEIN. Length = 218	gb AAC32842.	4788	2	823	43	59	HLDDDD01	pCMVSport 3.0
512	874459	(AF155575) peroxisomal D2,D4-dienoyl-CoA reductase [Mus musculus] >sp Q9WV68 Q9WV68 PEROXISOMAL D2,D4-DIENOYL-COA REDUCTASE. Length = 292	gb AAD38196. 1 AF1555	4789	_	531	85	06	HWLRA47	pSport1
513	874460			4790	47	244			HCRMX57	pSport1
514	874461			4791	285	488			HFPEC02	Uni-ZAP XR

HMEEI02 Lambda ZAP	pBluescript	pBluescript SK-	HUVGR86 Uni-ZAP XR	pBluescript SK-	pCMVSport 3.0	pBluescript SK-	pSport1	Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	Uni-ZAP XR	pSport1	HCQDD11 Lambda ZAP	pSport1
HMEEI02	HKCSZ54	H2CBM49	HUVGR86	HCYBN52	НDPFO58	H2CBC28	HCRQF18	HE2CI70	HSPAX64	HCRPE10	HTOJA79	HGBGI31	HCRMF12	нсбрр11	HCRPA46
		19	69							11				11	
		65	69							72				75	
283	743	262	1213	246	476	415	879	305	593	176	612	1150	320	187	325
32	480	185	2	10	186	146	724	36	360	39	355	938	3	2	101
4792	4793	4794	4795	4796	4797	4798	4799	4800	4801	4802	4803	4804	4805	4806	4807
		sp P39195 AL U8_HUMAN	dbj BAA09768 .1							sp P39194 AL U7_HUMAN				gb AAC51843.	
		III ALU SUBFAMILY SX WARNING ENTRY   III   sp P39195 AL   Length = 591	The KIAA0147 gene product is related to adenylyl cyclase. [Homo sapiens] >sp Q14160 Q14160 KIAA0147 PROTEIN (FRAGMENT). Length = 1551							ALU SUBFAMILY SQ WARNING ENTRY				(AF000996) ubiquitous TPR motif, Y isoform [Homo sapiens] Length = 1079	
874467	874468	874469	874470	874472	874473	874474	874475	874479	874480	874481	874482	874484	874485	874486	874492
515	516	517	518	519	520	521	522	523	524	525	526	527	528	529	530

pSport1	pSport1	Uni-ZAP XR	pBluescript	pCMVSport 2.0	pSport1	pCMVSport 3.0	HPMLY88 Uni-ZAP XR	pSport1	pSport1	pSport1
HCRPV94	HCRPX62	HFKIJ16	HL1SB93	HDTLA27	нснсл20	HLDOG81	HPMLY88	HIDAC50	HLYCA01	HCRNF16
94		86			22	100	66			82
93		98			40	100	66			75
562	615	649	417	403	1094	862	1213	199	271	463
2	235	2	1	41	8	1232	2	2	2	2
4808	4809	4810	4811	4812	4813	4814	4815	4816	4817	4818
emb CAA1974 2.1		gb AAF31436. 1 AF2163			gb AAC02603.	gb AAD34622. 1 AF1531	emb CAA3438 5.1			gb AAD16286. 1
(AL030998) dJ466I8.1 (Coagulation Factor V (Activated Protein C Cofactor), Coagulation Factor VIII (Procoagulant Component) and Ceruloplasmin (EC 1.16.3.1, Ferroxidase) LIKE) [Homo sapiens] >sp O75659 O75659 DJ466I8.1 (COAGULATION FACTOR V (ACTIVATED PROT		(AF216312) type II membrane serine protease [Homo sapiens] Length = 423			(AF045644) F57H12.7 gene product [Caenorhabditis gb AAC02603. elegans] >pir[T32979 T32979 hypothetical protein	(AF153191) nm23-H7 [Homo sapiens] >gb AAD34622.1 AF153191_1 (AF153191) nm23-H7 [Homo sapiens] >sp Q9Y5B8 Q9Y5B8 NM23-H7. Length = 376	54k protein (AA 1-504) [Canis familiaris] >emb[CAA60132.1  SRP 54 [Homo sapiens] >gb AAC50994.1  signal recognition particle [Homo sapiens] >pir S05197 S05197 signal recognition particle 54K protein - dog >pir S54143 S54143 SRP 54 protein - human (fragmen			(AF099066) serine/threonine-protein kinase NEK3 [Mus musculus] >sp Q9Z0X9 Q9Z0X9 SERINE/THREONINE-PROTEIN KINASE NEK3. Length = 509
874495	874498	874499	874503	874504	874505	874506	874508	874518	874519	874522
531	532	533	534	535	536	537	538	539	540	541

HOEKX93 Uni-ZAP XR	Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	pSport1	2 pSport1	pSport1	pSport1	pCMVSport 2.0
ноекх93	HTTFP72	HCRND05	HCRNP66	HAPCK19	HWLIN80	HWMBA02	HCRQI74	HCRMT48	HDTJ085
81	93	17	53			96	84	77	
75	93	64	32			96	83	77	
277	783	534	534	269	1036	920	572	235	826
68	_	-	-	150	830	3	192	2	962
4819	4820	4821	4822	4823		4825	4826	4827	4828
dbj BAA08226 .1	emb CAA0338 7.1	gb AAC16016. 1	emb CAB1428 8.1			gb AAC52071. 1	gb AAC26101. 1	dbj BAA31678 .1	
product specific to adipose tissue [Homo sapiens] >sp Q15847 Q15847 HYPOTHETICAL 7.9 KD PROTEIN. Length = 76	HUMAN NDR [unidentified] >emb CAA84485.1  Ndr protein kinase [Homo sapiens] >emb CAB39180.1  dJ108K11.2 (Ndr protein kinase) [Homo sapiens] >pir J38133 J38133 protein kinase (EC 2.7.1) Ndr - human >sp Q15208 Q15208 NDR PROTEIN KINASE. Length = 465	(AF062476) retinoic acid-responsive protein; STRA6 gb AAC16016. [Mus musculus] >sp O70491 O70491 RETINOIC 1 ACID-RESPONSIVE PROTEIN. Length = 670	similar to Na+/H+ antiporter [Bacillus subtilis] >dbj BAA12644.1  Yqkl [Bacillus subtilis] >pir B69967 B69967 Na+/H+ antiporter homolog yqkl - Bacillus subtilis >sp P54571 YQKI BACSU HYPOTHETICAL NA+/H+ ANTIPORTER IN ANSB-SPOIIM INTERGENIC REGION. Length			(AF022109) HsCdc18p [Homo sapiens] >gb AAB38317.1  Cdc6-related protein [Homo sapiens] >sp Q99741 Q99741 CDC6-RELATED PROTEIN. Length = 560	(AF034800) liprin-alpha3 [Homo sapiens] >sp AAC26101 AAC26101 Liprin-alpha3 (fragment). Length = 443	(AB014603) KIAA0703 protein [Homo sapiens] >sp O75185 O75185 KIAA0703 PROTEIN. Length = 963	
874524	874527	874528	874529	874531	874533	874534	874537	874540	874543
542	543	544	545	546	547	548	549	550	551

6	"	١
.,	ţ	J

Other	Uni-ZAP XR	pSport1	pSportl	pBluescript	HMIAD35 Uni-ZAP XR	pCMVSport 3.0	HDPAM86 pCMVSport 3.0	pSport1
HIBEM35	нЕ9QВ35	HCHMS55	HCRPG51	HKMLN95	HMIAD35	HSYAM68	HDPAM86	HNTMD17
	08	72	95	66		92	87	66
	75	72	95	66		91	87	66
1877	489	551	939	584	919	1140	1202	790
1452		8	_	24	623	238	138	2
4829	4830	4831	4832	4833	4834	4835	4836	4837
	emb CAA4044 5.1	emb CAA3887	gb AAB04028. 1	gb AAD44482. 1		gb AAB48435. 1	gb AAA58440.	dbj BAA91969 .1
	CHOX M product [Gallus gallus] >pir I50145 I50145 emb CAA4044 homeotic protein Hox M - chicken   5.1    5.1    5x P23459 HXD8 CHICK HOMEOBOX   PROTEIN HOX-D8 (CHOX-M). Length = 188	centromere autoantigen B (CENP-B) [Homo sapiens] emb CAA3887   >pir S18735 S18735 centromere protein B - human   >sp P07199 CENB_HUMAN MAJOR   CENTROMERE AUTOANTIGEN B   (CENTROMERE PROTEIN B) (CENP-B).   >gb AAB70165.1  (AF002714) centromere protein B; CENP-B [Homo sapiens] {	peroxisome proliferator activated receptor gamma 2 [Homo sapiens] >gb AAC51248.1  ligand activated transcription factor PPARgamma2 [Homo sapiens] >pir JC4859 JC4859 peroxisome proliferatoractivated receptor gamma-2 - human >sp Q15832 Q15832 PEROXISOME PR	(AF078850) steroid dehydrogenase homolog [Homo sapiens] >sp Q9Y6G8 Q9Y6G8 STEROID DEHYDROGENASE HOMOLOG. Length = 312		KHS1 [Homo sapiens] >sp Q9Y4K4 Q9Y4K4 KHS1. Length = 846	E2F-related transcription factor [Homo sapiens] >pir A48585 A48585 transcription factor DP-1 - human >sp Q14186 TDP1_HUMAN TRANSCRIPTION FACTOR DP-1 (E2F DIMERIZATION PARTNER 1) (DRTF1- POLYPEPTIDE-1) (DRTF1). Length = 410	(AK001902) unnamed protein product [Homo sapiens] Length = 484
874544	874545	874546	874550	874551	874552	874553	874556	874559
552	553	554	555	556	557	558	559	260

	874560	gap junction protein (aa 1-283) [Homo sapiens] >pir B29005 B29005 gap junction protein Cx32 - human >sp P08034 CXB1_HUMAN GAP JUNCTION BETA-1 PROTEIN (CONNEXIN 32) (CX32) (GAP JUNCTION 28 KD LIVER PROTEIN). Length = 283	emb CAA2785 6.1	4838	74	901	96	96	HEEAX65	HEEAX65 Uni-ZAP XR
	874561	Y SX WARNING ENTRY !!!	sp P39195 AL U8_HUMAN	4839	2	136	72	77	HHFJL44	Uni-ZAP XR
	874562	(AB018255) KIAA0712 protein [Homo sapiens] >sp O94820 O94820 KIAA0712 PROTEIN. Length = 1587	dbjBAA34432	4840	3	842	75	92	HWHGD94	pCMVSport 3.0
+	874563			4841	242	481			HWLAC81	pSport1
<del> </del>	874564	Nuclear localization signal at AA 569-573, 576-580, gb AAA58443. 579-583; acidic transcr. activ. domain 620-640;; lomeobox motif 653-676 [Homo sapiens] >pir A47456 A47456 down-regulated in adenoma (DRA) - human >sp P40879 DRA_HUMAN DRA PROTEIN (DOWN-REGULATED IN ADENO	gb AAA58443. 1	4842	109	363	98	91	нwгеоов	pSport1
+	874565			4843	583	828			HSQDM57	HSQDM57 Uni-ZAP XR
<del></del>	874567	STM-7 [Homo sapiens] >sp Q92749 Q92749 TYPE I emb CAA6322 PHOSPHATIDYLNOSITOL-4-PHOSPHATE 5- 4.1  KINASE BETA (EC 2.7.1.68) (STM-7 PROTEIN). >sb AAC50916.1  type I phosphatidylinositol-4-phosphate 5-kinase beta [Homo sapiens] {SUB 112-502} >sb AAC50914.1  type I phosphati	emb CAA6322 4.1	4844	2	1291	93	93	HTEJC93	Uni-ZAP XR
+	874569			4845	123	311			HWLMQ11	pSport1

HNSAD12 pSport1	HBJEN48 Uni-ZAP XR	56 HWMBM13 pSport1	H6BSM15 Uni-ZAP XR	HCQBD30 Lambda ZAP	HTEEZ83 Uni-ZAP XR	HBXCF35 ZAP Express	HWMBF85 pSport1	HCROA06 pSport1	HAPAY77 Uni-ZAP XR	HUSYW93 pSport1
PH 67	100 HB	MH 9	9H	HC	82 HT	EH —	HW	100 HC	HA	H
75 7	100	35 5			82 8		-	100		
485	527	539	468	428	1158	1154	277	426	937	618
102	м	E.	136	87	202	828	101	154	989	304
4846	4847	4848	4849	4850	4851	4852	4853	4854	4855	4856
emb CAB5599 0.1	emb CAA6201 3.1	gb AAA16358. 1			gb AAF19794. 1 AF1681			emb CAB7078		
(AL117555) hypothetical protein [Homo sapiens] >emb CAB55990.1  (AL117555) hypothetical protein [Homo sapiens] >pir T17300 T17300 hypothetical protein DKFZp564I1171.1 - human (fragment) >sp CAB55990 CAB55990 Hypothetical 14.8 kd protein (fragment). Length	NBK [Homo sapiens] >emb CAA18260.2  (AL022237) bK1191B2.2 (BCL2-interacting killer (apoptosis-inducing) (NBK, BP4, BIP1)) [Homo sapiens] >gb AAC50413.1  Bik [Homo sapiens] >gb AAC79124.1  apoptosis inducer Nbk [Homo sapiens] >spiens] >gb AAF01156.1  (AF174424) BCL	Eps8 [Mus musculus] >pir S39983 S39983 eps8 protein - mouse >sp Q08509 EPS8_MOUSE EPIDERMAL GROWTH FACTOR RECEPTOR KINASE SUBSTRATE EPS8. Length = 821			(AF168132) RU1 [Homo sapiens] >sp AAF19794 AAF19794 RU1. Length = 866			(AL137514) hypothetical protein [Homo sapiens] >emb CAB70780.1  (AL137514) hypothetical protein [Homo sapiens] >sp CAB70780 CAB70780 Hypothetical 11.2 kd protein. Length = 97		
874570	874571	874573	874577	874578	874580	874581	874584	874588	874590	874592
569	570	571	572	573	574	575	576	577	578	579

HCROE11 pSport1	HWLVF65 pSport1	HWLWU62 pSport1	HWLFG75 pSport1	HBCCB62 Uni-ZAP XR	HWLVN89 pSport1	HTXQF81 Uni-ZAP XR	HCQDD61 Lambda ZAP
82 H	H	H	64 H	92 H	H 16	H 100	田
79			38	92	06	001	
496	643	260	1139	1786	404	628	874
2	200	3	708	2	8	2	488
4857	4858	4859	4860	4861	4862	4863	4864
gb AAD13621. 1			gb AAC25392. 1	gb AAA51967.	emb CAB5602 6.1	gb AAA35499. 1	:
(AF123462) neurexin III [Homo sapiens]   >sp Q9Y486 Q9Y486 NEUREXIN III   (FRAGMENT). Length = 334			(AF070637) [Homo sapiens] >sp 075547 075547 HYPOTHETICAL 28.8 KD PROTEIN (FRAGMENT). Length = 256	carcinoembryonic antigen [Homo sapiens] >gb AAB59513.1  carcinoembryonic antigen precursor [Homo sapiens] >pir A36319 A36319 carcinoembryonic antigen precursor - human >sp P06731 CCEM_HUMAN CARCINOEMBRYONIC ANTIGEN PRECURSOR (CEA) (MECONIUM ANTIGEN 100) (	(AL117637) hypothetical protein [Homo sapiens] >emb CAB56026.1  (AL117637) hypothetical protein [Homo sapiens] >pir[T17336[T17336] hypothetical protein DKFZp4341225.1 - human (fragment) >sp CAB56026 CAB56026 Hypothetical 45.3 kd protein (fragment). Length	A4 protein [Homo sapiens] >gb AAB92356.1  A4 differentiation-dependent protein [Homo sapiens] >gb AAF05827.1 AF196779_4 (AF196779) A4 differentiation-dependent protein [Homo sapiens] >pir S32567 S32567 A4 protein - human >sp Q04941 A4P_HUMAN INTESTINAL ME	
874594	874595	874601	874603	874605	874607	874608	874609
280	581	582	583	584	585	586	587

Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1	Uni-ZAP XR	HOSOL09 Uni-ZAP XR	pSport1	pSport1	pBluescript SK-	pSport1	pSport1
HMCGZ52 Uni-ZAP XR	HDPMG95	HETAD58	HUFAT62	НОДСН47	HWLVI80	HNGBW96 Uni-ZAP XR	HOSOL09	HWLMK56	HWMBE67	H2CAA08	HCRNH24	HUFD017
99					70	95	70	98		86		
50					99	95	44	98		86		
1187	755	366	848	273	484	451	725	1130	468	432	1012	971
240	96	103	657	92	242	29	3	9	151	160	749	708
4865	4866	4867	4868	4869	4870	4871	4872	4873	4874	4875	4876	4877
emb CAA6987 5.1					dbj BAA91205	dbj BAA91013 .1	gb AAD32753. 1 AC0072	gb AAA35861. 1		gb AAB00195.		
UDP-GalNAc:polypeptide N-acetylgalactosaminyltransferase [Homo sapiens] >sp[000208 000208 UDP-GALNAC:POLYPEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE. Length = 578					(AK000496) unnamed protein product [Homo sapiens] Length = 239	d protein product [Homo 1	(AC007231) putative cation transport protein [Arabidopsis thaliana] >sp AAD32753 AAD32753 Putative cation transport protein. Length = 300	carcinoma-associated antigen GA733-2 [Homo sapiens] >gb AAB00775.1  carcinoma-associated antigen GA733-2 [Homo sapiens] >pir B48149 B48149 epithelial glycoprotein antigen GA733-2 precursor - human Length = 314		occludin [Homo sapiens] >gb AAC50451.1  occludin  gb AAB00195. [Homo sapiens] >pir G02533 G02533 occludin -   1  human >sp Q16625 OCLN_HUMAN OCCLUDIN. Length = 522		
874610	874611	874612	874614	874615	874618	874619	874620	874621	874622	874623	874624	874625
288	589	290	591	592	593	594	595	969	297	865	599	009

HE8QX06 Uni-ZAP XR	pSport1	pCMVSport 3.0	pSportl	pCMVSport 3.0	HNGGK17 Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1	pSport1
HE8QX06	HWMCF68	HWAGI58	HAAAA25	HHEIW79	HNGGK17	HCRQG35	HSODQ11	HWLMR54	HWLNI19
			100		97		97		96
			100		76		96		95
870	363	1434	099	365	2330	454	402	282	496
613	46	1213	58	186	120	236	175	1	392
4878	4879	4880	4881	4882	4883	4884	4885	4886	4887
			dbj BAA01374 .2		gb AAC34808. 1		.1		gb AAF07045. 1 AF1765
			p67 myc protein [Homo sapiens] >gb AAA20042.1  c-myc protein [Homo sapiens] {SUB 16-454} >emb CAA46984.1  this region determines c-myc mRNA stability [Homo sapiens] {SUB 395-454} Length = 454		(AF068229) lysyl hydroxylase 3 [Homo sapiens] >gb AAC39753.1  (AF046889) lysyl hydroxylase isoform 3 [Homo sapiens] >gb AAD45831.1 AC004876_4 (AC004876) lysyl hydroxylase 3 [Homo sapiens] >sp O60568 PLO3_HUMAN PROCOLLAGEN-LYSINE,2-OXOGLUTARATE 5-DIOXYGENA		(AB007917) KIAA0448 protein [Homo sapiens] >dbj BAA89250.1  (AB024568) heparan sulfate 2-sulfotransferase [Homo sapiens] >sp O75036 O75036 KIAA0448 PROTEIN. >sp BAA89250 BAA89250 Heparan sulfate 2-sulfotransferase. Length = 356		(AF176555) A-kinase anchoring protein 220 [Homo sapiens] >sp AAF07045 AAF07045 A-kinase anchoring protein 220. >dbj BAA31604.1  (AB014529) KIAA0629 protein [Homo sapiens] {SUB 1290-1901} Length = 1901
874626	874628	874630	874631	874632	874635	874636	874638	874639	874640
601	602	603	604	909	909	209	809	609	610

HFPHT42 Uni-ZAP XR	pCMVSport 3.0	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1.	pSport1	pSport1	pSport1	pSport1	pSport1	pSportl	pSport1	pSport1	Uni-ZAP XR
HFPHT42	HLWCT94	HWMBL25	HWLOU23	HWL0Z82	HWMBF50	HLYAZ23	HWLNL53	HWLOZ25	HWMBV27	HCRQH42	HWLOR14	HWMBB03	HWLOW57	HWL0077	HWLOZ54	HWLM019	HWLMA68	HWLNH87	HOOHE79	HWLOJ08	HBCBF08
73																		88			06
73																		79			06
328	879	532	464	479	<i>L</i> 69	390	133	366	625	470	201	260	651	227	318	310	467	703	513	208	009
23	673	308	237	162	128	199	444	154	320	312	64	27	809	96	193	2	234	47	208	35	1
4888	4889	4890	4891	4892	4893	4894	4895	4896	4897	4898	4899	4900	4901	4902	4903	4904		4906	4907	4908	4909
gb AAC50312.																		dbj BAA36338 .1			gb AAD48776. 1
interleukin-15 receptor alpha chain precursor [Homo  gb AAC50312. sapiens] >sp Q13261 Q13261 INTERLEUKIN-15  1  RECEPTOR ALPHA CHAIN PRECURSOR. >gb AAB88175.1  (AF035279) similar to interleukin-15 receptor alpha chain precursor [Homo sapiens] {SUB 37-267} Length = 267																		(AB015614) SET-binding protein (SEB) [Mus musculus] >sp Q9Z180 Q9Z180 SET-BINDING PROTEIN (SEB) (FRAGMENT). Length = 197			(AF151107) 3"-5" exonuclease TREX2 [Homo sapiens] >sp AAD48776 AAD48776 3"-5" exonuclease TREX2. Length = 236
874642	874644	874645	874646	874650	874651	874652	874653	874654	874655	874656	874657	874658	874659	874660	874662	874665	874667	874670	874671	874672	874673
611	612	613	614	615	919	617	618	619	620	621	622	623	624	625	979	627	879	629	630	631	632

633	874675			4910	458	655			HWHGZ23	pCMVSport 3.0
634	874678			4911	171	293			HWLOP85	pSport1
635	874679	(AL021918) b3418.1 (Kruppel related Zinc Finger protein 184) [Homo sapiens] >sp O60792 O60792 B3418.1 (KRUPPEL RELATED ZINC FINGER PROTEIN 184). Length = 751	emb CAA1727 8.1	4912	_	765	86	86	HUSGX66	pSport1
959	874680			4913	2	355			HCRQM95	pSport1
637	874682	Glucosamine-6-phosphate isomerase (EC 5.3.1.10) (glucosamine-6-phosphate deaminase). [Escherichia coli] >dbj[BAA35326.1  Glucosamine-6-phosphate isomerase (EC 5.3.1.10) (glucosamine-6- phosphate deaminase). [Escherichia coli] >gb AAA24191.1  glucosamine-	dbj BAA35321	4914	549	1106	06	06	HPWAI57	Uni-ZAP XR
638	874683			4915	2	424			НWLОQ35	pSport1
639	874684	G ENTRY !!!	sp P39189 AL U2_HUMAN	4916	-	150	89	73	HE2EA79	Uni-ZAP XR
640	874688	putative ATPase [Haematobia irritans] >splP46441 N2B HAEIR PUTATIVE ATPASE N2B (HFN2B). Length = 464	gb AAA91360. 1	4917	3	638	45	99	HWLOI43	pSport1
641	874689			4918	93	389			HCRQM44	pSport1
642	874695			4919	253	924			HCRMZ25	pSport1
643	874696			4920	1	246			HCROB95	pSport1
644	874697			4921	119	241			HWLXN82	pSport1
645	874699			4922	38	211			HWLXW08	pSport1
646	874700	(AF030881) pol polyprotein [Fugu rubripes] >sp AAC33526 AAC33526 Pol polyprotein (fragment). Length = 1187	gb AAC33526.	4923	71	349	32	53	HWLVR69	pSport1
647	874701	(AK000385) unnamed protein product [Homo sapiens] Length = 152	dbj BAA91131 .1	4924	1	129	78	81	H2CBD62	pBluescript SK-
648	874702			4925	89	397			HMSAQ57	Uni-ZAP XR
649	874703			4926	403	829			HCROD17	pSport1

pBluescript SK-	pBluescript SK-	Uni-ZAP XR	pBluescript SK-	pBluescript SK-	pBluescript SK-	Uni-ZAP XR	Uni-ZAP XR	Uni-ZAP XR	HFRAM50 Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR
H2CBN90	H2CBP17	HTTDU01	Н2СВН38	H2CBX48	H2CBT32	HAGBH67	HE2LX05	HAHCU44	HFRAM50	HAJBD60	НТРНК <i>47</i>
83	100	25					100	91	08	99	86
78	100	94					95	16	73	48	86
821	539	686	473	363	1141	1105	200	876	145	811	1341
ε	09	81	228	-	926	260	339	-	303	=	
4927	4928	4929	4930	4931	4932	4933	4934	4935	4936	4937	4938
dbj BAA34216 .1	emb CAB7542 7.1	dbj BAA91645 .1					gb AAB53629.	dbj BAA90980 .1	gb AAB61715. 1	gb AAA84746. 1	emb CAB4322 9.1
(AB005549) atypical PKC specific binding protein [Rattus norvegicus] >pir[T13948 T13948 atypical protein kinase C isotype-specific interacting protein ASIP - rat >sp Q9Z340 Q9Z340 ATYPICAL PKC SPECIFIC BINDING PROTEIN. Length = 1337	(AJ271784) chromokinesin [Homo sapiens] Length = emb CAB7542 1232	(AK001355) unnamed protein product [Homo sapiens] Length = 291					beta-galactosidase [Expression vector pBSII- LUCINT] Length = 69	protein product [Homo	to sapiens] NRF2: FUNCTION . Length =	POM1 [Plasmodium chabaudi chabaudi] >sp Q25658 Q25658 POM1 (FRAGMENT). Length = 597	(AL049989) hypothetical protein [Homo sapiens] >emb CAB43229.1  (AL049989) hypothetical protein [Homo sapiens] >pir T08691 T08691 hypothetical protein DKFZp564F052.1 - human (fragment) >sp Q9Y3Z7 Q9Y3Z7 HYPOTHETICAL 48.5 KD PROTEIN (FRAGMENT). Length = 42
874704	874707	874708	874709	874710	874711	874713	874714	874715	874717	874718	874719
059	651	652	653	654	655	959	657	859	659	099	661

662	874720	RLF [Homo sapiens] >sp[Q13129]Q13129 ZN-15 gb RELATED ZINC FINGER PROTEIN (RLF). 11 Length = 1914	gb AAC50396. 1	4939	1	696	36	49	HAMGM27	49 HAMGM27 pCMVSport 3.0
663	874723			4940	25	129			HWLXA56	pSport1
664	874724	CCHA PROTEIN PRECURSOR. [Escherichia coli] db >gb AAC75510.1  (AE000332) detox protein [Escherichia coli] >pir H65020 H65020 cchA protein - Escherichia coli (strain K-12) >sp BAA16335 BAA16335 CCHA PROTEIN PRECURSOR >sp P77606 EUTM_ECOLI	dbj BAA16335 .1	4941	176	К	86	86	HBGMC86	HBGMC86 Uni-ZAP XR
999	874725			4942	146	289			HOSPA23	Uni-ZAP XR
999	874726			4943	473	712			HBAHC42	pSport1
299	874727			4944	899	793			HUSGQ45	pSport1
899	874728			4945	396	979			HBMXP34	HBMXP34   Uni-ZAP XR
699	874732			4946	1300	1635			HHEME74	pCMVSport 3.0
029	874737			4947	81	281			HCNDN66	HCNDN66 Lambda ZAP
671	874741	alanine aminotransferase [Homo sapiens] Length = db 493	dbj BAA01186 .1	4948	12	842	99	82	H2CBI61	pBluescript SK-
672	874744			4949	111	329			HCQAE09	HCQAE09 Lambda ZAP
673	874745	IIII ALU SUBFAMILY SQ WARNING ENTRY   IIII   sp P39194 AL   Length = 593	sp P39194 AL U7_HUMAN	4950	259	357	75	82	HCNDP23	Lambda ZAP II
674	874746			4951	1	225			нсове66	HCQBE66 Lambda ZAP
675	874747			4952	68	178	.,		нсоак59	Lambda ZAP II

HCQAR64 Lambda ZAP	pSport1	Lambda ZAP II	pSport1	pSport1	Lambda ZAP II	HCQBH60 Lambda ZAP	pSport1	Lambda ZAP II	HCQCC66 Lambda ZAP	Uni-ZAP XR	HCQCB62 Lambda ZAP	HCQCC13 Lambda ZAP	HCQCF83 Lambda ZAP	HCQAF27 Lambda ZAP	HCQCJS6 Lambda ZAP	HCQCD88 Lambda ZAP
HCQAR64	HWMAC48	нсове76	HWLCA32	HWLHH20	нсов172	нсовн60	HHMMB17	<b>НСQСВ28</b>	9922ОЭН	HOELS72	нсосв62	нсосс13	нсосғ83	HCQAF27	нсосл56	НСОСD88
96	29															
96	55															
248	362	333	261	354	492	478	166	442	282	225	125	153	199	171	1132	147
m	<sub>60</sub>	202	58	22	307	347	2	233	-	73	<sub>6</sub>	-	182	-	929	-
4953	4954	4955	4956	4957	4958	4959	4960	4961	4962	4963	4964	4965	4966	4967	4968	4969
gb AAA58477.	emb CAB6919 5.1															
fork head-related protein [Homo sapiens] >sp[P55318]HN3G_HUMAN HEPATOCYTE NUCLEAR FACTOR 3-GAMMA (HNF-3G) (FORK HEAD-RELATED PROTEIN FKH H3). Length = 347	unnamed protein product [unidentified] Length = 180 emb CAB6919 5.1															
874748	874749	874750	874751	874752	874753	874754	874755	874756	874757	874758	874759	874760	874763	874764	874765	874766
929	119	829	629	089	681	682	683	684	685	989	289	889	689	069	691	692

CTP synthetase homolog [Mus musculus] >sp P70303 P70303 CTP SYNTHETASE HOMOLOG (CTPSH). Length = 586		gb AAB17729. 1	4970	2	598	70	78	HE8OJ09	HE8OJ09 Uni-ZAP XR
			4971	1	219			HCQCR67	HCQCR67   Lambda ZAP II
120001	expressed ubiquitously with strong expression in brain [Homo sapiens] >gb AAD15417.1  (AC004912) KIAA0193 [Homo sapiens] >sp Q12765 Y193_HUMAN HYPOTHETICAL PROTEIN KIAA0193. >sp AAD15417 AAD15417 KIAA0193. Length = 346	dbj BAA12106 .1	4972	477	1379	47	89	нРна <i>а</i> 27	HPHAA27 Uni-ZAP XR
1			4973	209	406			HCROV23	pSport1
l			4974	287	490			HCRMZ75	pSport1
	:		4975	171	557			HCRMZ85	pSport1
			4976	1	162			HCROM08	pSport1
			4977	312	545			HBIPL82	pCMVSport 3.0
	protein product [Homo	dbj BAA90891 .1	4978	539	961	92	92	HBXBV89	HBXBV89 ZAP Express
	.1H 3" region) - human Length pir B34087 B3 4087	pir B34087 B3 4087	4979	14	250	51	99	HCRPM45	
			4980	1	168			нсост75	Lambda ZAP II
			4981	1	237			HCRP092	pSport1
			4982	3	314			HCRNM87	pSport1
			4983	448	642			HBJFU36	Uni-ZAP XR
			4984	43	330			HCRPZ29	pSport1
			4985	2	232			HCRON58	pSport1

pSport1	ambda ZAP II	pBluescript SK-	pBluescript SK-	pSport1	pSport1	pBluescript SK-	pSport1	pBluescript SK-	pSport1	pSport1	pBluescript SK-	pSport1
HCRNG90	HCQDT67 Lambda ZAP	HCYAC32	нсувк32	HWMCE07	HCROL83	нсувм89	HCRNX33	HCYBM31	HDAAX73	HDACJ67	H2CBL90	HPC0E53
68		86	87			87						
89		86	81			87					•	
947	388	909	127	291	362	522	388	669	415	469	471	655
87	146	249	=	1	123	112	53	505	311	353		524
4986	4987	4988	4989	4990	4991	4992	4993	4994	4995	4996	4997	4998
emb CAA3927		dbj BAA76932 .1	emb CAA0647			gb AAB62941. 1						
5"-nucleotidase [Homo sapiens] >pir S11032 S11032   emb CAA3927   5"-nucleotidase (EC 3.1.3.5) precursor - human   >sp P21589 SNTD_HUMAN 5"-NUCLEOTIDASE   PRECURSOR (EC 3.1.3.5) (ECTO-NUCLEOTIDASE) (5"-NT) (CD73 ANTIGEN).   >gb AAA96950.1 5"-nucleotidase [Homo sapiens] {		(AB024705) fls485 [Homo sapiens] >sp Q9Y2M2 Q9Y2M2 FLS485. Length = 353	(AJ005324) glutamate permease [synthetic construct] emb CAA0647 > emb CAA06474.1  (AJ005327) glutamate permease [1.1  [synthetic construct] > emb CAA06477.1  (AJ005330) glutamate permease [synthetic construct] > gb AAA24514.1  gltS [Escherichia coli] {SUB 437-459} Length = 45			(AF007551) Bet1p homolog [Homo sapiens] >gb AAD47132.1 AC006378_1 (AC006378) Bet1p homolog [Homo sapiens] >sp O15155 O15155 BET1P HOMOLOG. >sp AAD47132 AAD47132 Bet1p homolog. Length = 118						
874787	874788	874790	874791	874793	874795	874796	874797	874800	874801	874802	874803	874804
709	710	711	712	713	714	715	716	717	718	719	720	721

pCMVSport 3.0	pSport1	pCMVSport 2.0	pBluescript SK-	pCMVSport 2.0	Uni-ZAP XR	Uni-ZAP XR	pCMVSport 2.0	pCMVSport 3.0	HE2LN12 Uni-ZAP XR	pSport1	Uni-ZAP XR	Uni-ZAP XR	Uni-ZAP XR
HDPGS84	HCRMQ21	HDTBM35	HCYBL83	HDTJE91	HE6BJ48	HE8NK63	HDTHF30	HDPRY54	HE2LN12	HWLUR88	HE8SB04	НЕ9QМ31	HTELU32
						51							74
						33							73
319	236	357	359	220	605	843	752	926	761	326	1108	392	270
188	123	256	3	2	306	-	618	672	516	78	881	108	
4999	2000	5001	5002	5003	5004	5005	2006	5007	2008	5009	5010	5011	5012
						emb CAA9812 0.1							emb CAB5602 1.1
						cDNA EST EMBL:M88866 comes from this gene [Caenorhabditis elegans] >pir T20358 T20358 hypothetical protein D2030.8 - Caenorhabditis elegans >sp P90793 P90793 D2030.8 PROTEIN. Length = 648							(AL117629) hypothetical protein [Homo sapiens] >emb CAB56021.1  (AL117629) hypothetical protein [Homo sapiens] >pir[T17331[T17331] hypothetical protein DKFZp434C245.1 - human (fragment) >sp CAB56021 CAB56021 Hypothetical 21.0 kd protein (fragment). Length
874805	874807	874809	874810	874812	874813	874815	874816	874818	874819	874820	874821	874822	874827
722	723	724	725	726	727	728	729	730	731	732	733	734	735

HEMGV90 Uni-ZAP XR	pCMVSport 2.0	pSport1	pSport1	Uni-ZAP XR	pSport1	HHGDC54   Lambda ZAP	HMSCD54 Uni-ZAP XR	pSport1	HHGDL18 Lambda ZAP	HOSMQ26 Uni-ZAP XR	pSport1
HEMGV90	HDTMC78	HFOXN77	HWLMW61	HHFLR55	HWLQ014	HHGDC54	HMSCD54	HISCH48	HHGDL18	HOSMQ26	HISDK89
57			06	94							
47			06	94							
536	202	774	155	1000	551	250	719	575	424	424	130
r.	11	505	E.	2	249	95	295	339	188	2	41
5013	5014	5015	5016	5017	5018	5019	5020	5021	5022	5023	5024
emb CAB0503 0.1			gb AAA81646. 1	gb AAA64911. 1							
predicted using Genefinder; Similarity to E.coli guanosine-3", 5"-bis(diphosphate)-pyrophosphohydrolase (SW:P17580) [Caenorhabditis elegans] >pir[T28096[T28096 hypothetical protein ZK909.3 - Caenorhabditis elegans >sp O18307 O18307 ZK909.3 PROTEIN. Leng			methylthioadenosine phosphorylase [Homo sapiens] >pir[138969 138969 5"-methylthioadenosine phosphorylase (EC 2.4.2.8) - human >sp Q13126 MTAP_HUMAN 5"-METHYLTHIOADENOSINE PHOSPHORYLASE (EC 2.4.2.28) (MTA PHOSPHORYLASE) (MTAPASE). Length = 283	Gem [Homo sapiens] >pir A54575 A54575 35K GTP-binding protein Gem - human >sp P55040 GEM_HUMAN GTP-BINDING PROTEIN GEM (GTP-BINDING MITOGEN- INDUCED T-CELL PROTEIN) (RAS-LIKE PROTEIN KIR). Length = 296							
874828	874829	874830	874832	874835	874836	874837	874843	874844	874845	874847	874849
736	737	738	739	740	741	742	743	744	745	746	747

pSport1	pSport1	pSport1	Uni-ZAP XR	HMAMA02 Uni-ZAP XR	pCMVSport 2.0	pSport1	pCMVSport 2.0	pCMVSport 3.0	ZAP Express	pBluescript	pSport1	pCMVSport 2.0	pCMVSport 3.0	pCMVSport 2.0
HLSAA22	HFOXR45	HWLOV52	HKCAA14	НМАМА02	HKABV02	HKGBD56	HKACE03	HBIOR20	HKEAA44	HKLSA63	HKGC122	HOGDO85	HLDOX53	НКАНЈ56
96				91				50						80
95				06				31						80
784	178	589	488	_	402	262	427	191	236	335	619	1062	528	859
62	23	2	261	390	7	59	2	6	3	222	491	962	1	38
5025	5026	5027	5028	5029	5030	5031	5032	5033	5034	5035	5036	5037	5038	5039
emb CAA8054				gb AAA67566. 1				dbj BAA90265 .1						gb AAD34079. 1 AF1518
M130 antigen [Homo sapiens] >emb CAB45233.1  CD163 [Homo sapiens] >pir I38003 S36077 M130 antigen - human >sp Q07898 Q07898 M130 ANTIGEN PRECURSOR. Length = 1116				f198 [Escherichia coli] >gb AAC77483.1  (AE000453) orf, hypothetical protein [Escherichia coli] >pir E65179 E65179 hypothetical 22.4 kD protein in trpT-pssR intergenic region - Escherichia coli (strain K-12) >sp P22788 YIFA_ECOLI HYPOTHETICAL 22.4 KD PROT				(AB022914) TES101RP [Mus musculus] >sp BAA90265 BAA90265 TES101RP. Length = 250						(AF151842) CGI-84 protein [Homo sapiens] >sp AAD34079 AAD34079 CGI-84 protein. Length = 213
874851	874852	874854	874855	874856	874857	874858	874859	874864	874865	874866	874867	874870	874871	874873
748	749	750	751	752	753	754	755	756	757	758	759	092	761	762

	8/48/5	ORF4 [Rattus norvegicus] >pir S21348 S21348 probable pol polyprotein-related protein 4 - rat >sp Q63306 Q63306 LONG INTERSPERSED REPETITIVE DNA CONTAINING 7 ORF"S. Length = 275	emb CAA3764 7.1	5040	376	444	52	72	HLTBL32	Uni-ZAP XR
	874876			5041	250	468			HLTHZ36	HLTHZ36 Uni-ZAP XR
765	874877			5042	48	158			HMEES39	Lambda ZAP II
99/	874879			5043	118	378			HMKA091	pSport1
-	874880			5044	317	889			HLYAQ21	pSport1
	874881	(AF022857) neuropilin-2(b0) [Mus musculus] >sp 035376 035376 NEUROPILIN 2. Length = 901	gb AAC53380.	5045	74	490	82	84	HCRNL20	pSport1
<del>                                     </del>	874885	GTP-binding protein - mouse Length = 198	pir S39543 S39 543	5046	128	730	66	100	HSYDX40	pCMVSport 3.0
770	874886	(AK000178) unnamed protein product [Homo sapiens] Length = 358	dbj BAA90992 .1	5047	2	913	90	90	НWLОQ11	pSport1
	874888	(AF129534) F-box protein Fbx4 [Homo sapiens] >sp AAF04468 AAF04468 F-box protein Fbx4. Length = 387	gb AAF04468. 1 AF1295	5048	409	1170	82	84	HMTAD91	pCMVSport 3.0
772	874889			5049	2	136			HOSFI36	Uni-ZAP XR
773	874890			5050	998	849			HHEYM94	pCMVSport 3.0
774	874891	SPIN protein [Homo sapiens] >gb AAC08315.1  (AF038969) general transcription factor 2-1 [Homo sapiens] >sp O15359 O15359 SPIN PROTEIN. Length = 957	emb CAA7516 3.1	5051	2	601	28	88	HPWCL64	HPWCL64   Uni-ZAP XR
-	874892			5052	2	538			HNTSQ62	pSport1
	874893	ROK-ALPHA. Length = 1379	gb AAB37540. 1	5053	177	395	<i>L</i> 9	71	HRDDU54	Uni-ZAP XR
-	874894			5054	-	180			HRDBA25	Uni-ZAP XR

HSRAJ45 Uni-ZAP XR	pBluescript SK-	pSport1	pSport1	HRDFM44 Uni-ZAP XR	pBluescript SK-	HSUBX76 Uni-ZAP XR	HNEAF57 Uni-ZAP XR	pSport1	pBluescript
HSRAJ45	HSABG91	HWLGN30	HSPAL74	HRDFM44	HCYBJ79		HNEAF57	HWLRA09	HSUSB86
					98	100	45		
					74	100	24		
758	192	1191	202	346	147	341	1044	317	941
543	338	880	2	149	7	m	184	135	552
5055	5056	5057	5058	5059	2060	5061	5062	5063	5064
					gb AAC51322.	emb CAA3870	emb CAA7961		
					sin3 associated polypeptide p18 [Homo sapiens] >gb AAD41090.1 AF153608_1 (AF153608) sin3 associated polypeptide [Homo sapiens] >sp O00422 SP18_HUMAN SIN3 ASSOCIATED POLYPEPTIDE P18. Length = 153	Cks1 protein homologue [Homo sapiens] >pit B36670 B36670 cell division control protein CKS2 - human >sp P33552 CKS2_HUMAN CYCLIN-DEPENDENT KINASES REGULATORY SUBUNIT 2 (CKS-2). Length = 79	cDNA EST yk425a6.3 comes from this gene; cDNA EST yk406e6.3 comes from this gene; cDNA EST yk425a6.5 comes from this gene; cDNA EST yk480c6.5 comes from this gene; cDNA EST yk406e6.5 comes from this gene [Caenorhabditis elegans] >sp CAA79619 CAA79619 F02A		
874895	874896	874897	874898	874899	874900	874902	874903	874904	874905
778	779	780	781	782	783	784	785	982	787

788	874906	bumetanide-sensitive Na-K-Cl cotransporter [Homo sapiens] >pir A57187 A57187 bumetanide-sensitive Na-K-Cl cotransporter - human >spp55011 NKC1_HUMAN BUMETANIDE-SENSITIVE SODIUM-(POTASSIUM)-CHLORIDE COTRANSPORTER 1 (BASOLATERAL NA-K-CL SYMPORTER). Length	gb AAC50561. 1	5065	2	1027	66	66	HOSAK80	HOSAK80 Uni-ZAP XR
789	874907	(AF049089) casein kinase I gamma 3 [Homo gb AAD26 sapiens] >sp Q9Y6M4 Q9Y6M4 CASEIN KINASE I 1 AF0490 GAMMA 3. Length = 447	gb AAD26525. 1 AF0490	2066	2	196	77	77	HE8TM43	HE8TM43 Uni-ZAP XR
790	874908	oiens] >pir PC4035 PC4035 50K nuclear protein - human 1 Q13171 AH ANTIGEN h = 1017	gb AAA86889. 1	5067	2	235	71	77	HTTBS45	Uni-ZAP XR
791	874909			5068	24	599			HLYAI14	pSport1
792	874912			6905	289	405			HODFU18	HODFU18 Uni-ZAP XR
793	874914			5070	22	459			HTXCZ25	HTXCZ25 Uni-ZAP XR
794	874917			5071	171	548			HWDAU63	HWDAU63 pCMVSport 3.0
795	874924	homeotic protein Hox-3.6 - mouse >sp P31257 HXCA_MOUSE HOMEOBOX PROTEIN HOX-C10 (HOX-3.6) >pir B60941 B60941 homeotic protein hox 3I - human (fragment) {SUB 268-333} >sp P31312 HXCB_MOUSE HOMEOBOX PROTEIN HOX-C11 (HOX-3.7) (FRAGMENT). {SUB 268-327} >gb A	pir A56552 A5 6552	5072	33	647	88	92	НWННG74	HWHHG74 pCMVSport 3.0
796	874925			5073	156	407			HWLIE53	pSport1

78 HWLLR30 pSport1	62 HLYCA86 pSport1	HDPTI77 pCMVSport	HWBDT18 pCMVSport 3.0	51 HWLMV62 pSport1	100 H2MAC06 pBluescript SK-	HTNAL08 pBluescript SK-	HCQAM40 Lambda ZAP	HWLQA72 pSport1
78 7	52 6	-		31 5	100 10			
183	173	814	1050	463	763 1	547	341	282
-	3	161	874	2	2	374	114	85
5074	5075	5076	5077	5078	5079	2080	5081	5082
gb AAB40602. 1	pir B40201 B4 0201			emb CAB7084 0.1	emb CAA5812 7.1			
homeobox protein Cdx1 [Homo sapiens] >sp P47902 CDX1_HUMAN HOMEOBOX PROTEIN CDX-1 (CAUDAL-TYPE HOMEOBOX PROTEIN 1). >gb AAC50237.1  caudal-type homeobox protein [Homo sapiens] {SUB 63-265} >gb AAA80284.1  caudal-type homeobox protein [Homo sapiens] {SUB 1	artifact-warning sequence (translated ALU class B) - $pir B40201 B4$ human Length = $301$			(AL137608) hypothetical protein [Homo sapiens] >emb CAB70840.1  (AL137608) hypothetical protein [Homo sapiens] >sp CAB70840 CAB70840 Hypothetical 46.0 kd protein (fragment). Length = 398	neutrophil gelatinase associated lipocalin [Homo sapiens] >sp P80188 NGAL_HUMAN NEUTROPHIL GELATINASE-ASSOCIATED LIPOCALIN PRECURSOR (NGAL) (P25) (25 KD ALPHA-2-MICROGLOBULIN-RELATED SUBUNIT OF MMP-9) (LIPOCALIN-2) (ONCOGENE 24P3). >gb AAD14168.1 S75256_1			
874926	874927	874928	874929	874930	874931	874932	874933	874934
797	798	799	800	801	802	803	804	805

pBluescript SK-	Uni-ZAP XR	pSport1	pSport1	pSport1	pSport1	pCMVSport 2.0	pSport1	HUVFU42 Uni-ZAP XR	pCMVSport 2.0	pSport1	pSport1
H2LAD85	HFKHN59	HWLRB64	HWLQB30	HWLRS70	HWLR068	HDLAZ62	HCRPS91	HUVFU42	HDTAC50	HWLWO06	HWLWP88
93			93				51	93	88		
93			93				31	93	88		
1085	594	619	63	132	717	373	941	1607	1358	423	200
<i>E</i>	388	320	-	-	1	2	294	m	E.	94	3
5083	5084	5085	9809	5087	5088	5089	2090	5091	5092	5093	5094
emb CAA4506			sp P39194 AL U7_HUMAN				gb AAD34068. 1 AF1518	gb AAA52412. 1	gb AAA59461.		
TFIIE-beta [Homo sapiens] >gb AAB20414.1  emb CAA4506 general transcription factor IIE 34 kda subunit, TFIIE 9.1  34 kda subunit [human, Peptide, 291 aa] [Homo sapiens] >pir S29292 S29292 transcription factor TFIIE-beta - human >sp P29084 T2EB_HUMAN TRANSCRIPTION INITIATI			IIII ALU SUBFAMILY SQ WARNING ENTRY IIII spp3194 AL Length = 593				(AF151831) CGI-73 protein [Homo sapiens] >sp Q9Y382 Q9Y382 CGI-73 PROTEIN. Length = 335	ets2 protein [Homo sapiens] >gb AAB94057.1  (AF017257) erythroblastosis virus oncogene homolog 2 protein [Homo sapiens] >pir B32066 TVHUE2 transcription factor ets-2 - human >sp P15036 ETS2_HUMAN C-ETS-2 PROTEIN. >gb AAA52411.1  ets protein [Homo sapiens]	keratin 18 [Homo sapiens] >gb AAA59463.1  keratin gb AAA59461.  18 precursor [Homo sapiens] >emb CAA31375.1  cytokeratin 18 [Homo sapiens] >pir S05481 S05481 keratin 18, type 1, cytoskeletal - human >sp D05783 K1CR_HUMAN KERATIN, TYPE I CYTOSKELETAL 18 (CYTOKERATIN 18)		
874936	874937	874938	874939	874944	874946	874951	874957	874958	874962	874965	874970
908	807	808	608	810	811	812	813	814	815	816	817

818	874972	tumor-associated antigen [Homo sapiens] >pir A36056 A36056 tumor-associated antigen CO-029 - human >sp P19075 C002_HUMAN TUMOR-ASSOCIATED ANTIGEN CO-029. Length = 237	gb AAA35709.	5095	207	929	46	94	HWLHW19	pSport1
819	874973			9605	641	883			HNTA183	pCMVSport 3.0
820	874974			5097	2	310			HWLWS24	pSport1
821	874975			8609	3	431			HWLWP62	pSport1
822	874976	IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	sp Q9Y6Y5 Q9 Y6Y5	5099	_	363	81	87	HOENV16	Uni-ZAP XR
823	874977			5100	19	186			HCRPM57	pSport1
824	874978			5101	84	227			HWLQT35	pSport1
825	874979			5102	689	823			HTWBQ51	pSport1
826	874980			5103	39	500			HWLWS65	pSport1
827	874981			5104	∞	175			HCRQC24	pSport1
828	874983	precursor polypeptide (AA -21 to 782) [Homo sapiens] >pir A35954 A35954 endoplasmin precursor 1.1  - human >sp P14625 ENPL_HUMAN ENDOPLASMIN PRECURSOR (94 KD GLUCOSE-REGULATED PROTEIN) (GRP94) (GP96 HOMOLOG) (TUMOR REJECTION ANTIGEN 1). Length = 803	emb CAA3326 1.1	5105	-	1110	93	93	HTFNM11	pSport1
829	874984			5106	748	939			HFTUG95	pSport1
830	874985			5107	617	808			HSRFC02	Uni-ZAP XR
831	874989			5108	155	448			HCRPC43	pSport1
832	874990	IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593	sp P39194 AL U7_HUMAN	5109	318	362	89	75	HMSPB24	Uni-ZAP XR
833	874991	<pre>fomo sapiens] 168_HUMAN HYPOTHETICAL IAA0168. Length = 326</pre>	dbj BAA11485  -11	5110	3	434	40	57	HWLW183	pSport1

HCQBI18 Lambda ZAP	E49 pSport1	[59 pSport]	86 pSport1	130 pSport1	[54 pSport]	69 pSport1	K08 pSport1	130 Lambda ZAP II	V59 pSport1	118 pSport1	101 pSport1	-	43 Uni-ZAP XR	346 pSport1	SS7 nSnort1
нсові	HWMBE49	HCRPH59	HCRPJ86	HCRPH30	HCRPH54	HCRPH69	HWLVX08	HKLAA30	HWLVW59	HWLJN18	HCROH01	HCRPJ81	HETGS43	HWLRS46	HWI.RS57
	92	78	66		77		89	77	_	08		_		66	
	8	75	86	_	89		45	8		77				66	
412	713	449	666	146	378	347	377	222	528	569	219	260	423	552	252
11	438	369	-	3	259	252	330	64	346	m	3	147	211	-	-
5111	5112	5113	5114	5115	5116	5117	5118	5119	5120	5121	5122	5123	5124	5125	5126
	dbj BAA03369 .1	sp P39192 AL US_HUMAN	dbj BAA91761 .1		sp P39193 AL U6_HUMAN		pir A25313 GN HUL1	dbj BAA81747  -1		pir A49963 A4 3932				gb AAA53500. 1	
	calpain [Rattus norvegicus] Length = 703	IIII ALU SUBFAMILY SC WARNING ENTRY IIII sp P39192 AL Length = 585	(AK001568) unnamed protein product [Homo sapiens] Length = 718		!!!! ALU SUBFAMILY SP WARNING ENTRY !!!! sp P39193 AL Length = 593		retrovirus-related reverse transcriptase pseudogene - human >sp P08547 LIN1_HUMAN LINE-1 REVERSE TRANSCRIPTASE HOMOLOG. Length = 1259	(AB002631) collectin 34 [Homo sapiens]   >sp Q9Y6Z7 Q9Y6Z7 COLLECTIN 34. Length = 277		mucin 2 precursor, intestinal - human (fragments) pir A49963 A4				cytochrome P450 IID6 [Homo sapiens] Length = 497 [gb AAA53500.	
874992	874993	874994	874995	874996	874997	874998	874999	875001	875002	875003	875004	875005	875007	875008	875009
834	835	836	837	838	839	840	841	842	843	844	845	846	847	848	840

[			<u>-</u>	-			امر	T	r		I~				ı —	0.			
pSport1	pSport1	pSport1	Uni-ZAP XR	pSport1	pSport1	pSport1	Uni-ZAP XR	pSport1	pSport1	pSport1	Uni-ZAP XR	pSport1	pSport1	pSport1	pSport1	Lambda ZAP II	pSport1	pSport1	pSport1
HUSJO81	HCRPF66	HRMAF73	HMSMR90	99МОТМН	HCRON47	HWLRV45	HFGAB06	HWLVA35	HCRPQ86	HCROZ20	HFPKD18	HCROS59	HCROR65	HZAAD77	HCRPA12	HMEKZ86	HCRPR27	HCRPQ46	HCRPN09
86								84	97	93	100								
86								82	76	68	100								
373	247	909	185	454	331	259	1105	304	210	380	242	261	226	142	303	1678	995	312	236
2	62	460	3	281	146	50	818	128		147	99	94	2	44	178	1325	357	184	75
5127	5128	5129	5130	5131	5132	5133	5134	5135	5136	5137	5138	5139	5140	5141	5142	5143	5144	5145	5146
gb AAA60235.								sp P39192 AL US_HUMAN	dbj BAA31603 .1	sp P39193 AL U6_HUMAN	gb AAD27738. 1 AF1329								
ubiquinone-binding protein (QP) [Homo sapiens] >gb AAA60238.1  ubiquinone-binding protein precursor [Homo sapiens] >emb CAA31926.1  ubiquinone-binding protein (AA 1 - 111) [Homo sapiens] >pir A32450 A32450 ubiquinone-binding protein QP-C - human >sp P1492								IIII ALU SUBFAMILY SC WARNING ENTRY IIII sp P39192 AL Length = 585	KIAA0628 protein [Homo sapiens] 075123 KIAA0628 PROTEIN. Length	iiii ALU SUBFAMILY SP WARNING ENTRY iiii sp P39193 AL Length = 593	(AF132963) CGI-29 protein [Homo sapiens] >sp Q9Y318 Q9Y318 CGI-29 PROTEIN. Length = 242								
875011	875017	875018	875019	875020	875024	875025	875027	875029	875032	875034	875035	875036	875037	875038	875042	875044	875045	875046	875047
850	851	852	853	854	855	958	857	858	859	098	861	862	863	864	865	998	298	898	698

875048 875049				5147	35	205			HCRPK03 HWLHY62	pSport1 pSport1
875053				5149	81	509			H2CBP44	pBluescript SK-
875055				5150	3	242			HCROW75	pSport1
875056				5151	1	87		-	HCROW65	pSport1
875058 (AK000361) unnamed protein product [Homo sapiens] Length = 692			dbj BAA91111	5152	18	260	66	66	HPJCF45	Uni-ZAP XR
875059				5153	170	295			HCRON87	pSport1
875060				5154	240	632			HIBEL82	Other
875061				5155	41	235			HCRPE83	pSport1
875062 (AJ009937) nuclear hormone receptor PRR2-A [Homo sapiens] >splCAB55492 CAB55492 Nuclear hormone receptor PRR2-A. >emb CAB55494.1  (AJ009937) nuclear hormone receptor PRR2-B [Homo sapiens] {SUB 56-397} Length = 397	(AJ009937) nuclear hormone recepto [Homo sapiens] >sp CAB55492 CAB hormone receptor PRR2-A. >emb CA (AJ009937) nuclear hormone recepto [Homo sapiens] {SUB 56-397} Lengt	clear	emb CAB5549 2.1	5156	rs.	95	87	06	н <b>w</b> LuQ22	pSport1
875063 homeobox protein Gsh-1 [Mus musculus] >pir S63604 S63604 homeobox protein Gsh-1 - mouse >sp P31315 GSHI_MOUSE HOMEOBOX PROTEIN GSH-1. >pir A37290 A37290 homeotic protein Gsh-1 - mouse (fragment) {SUB 146-205} >sp G601819 G601819 HPX HOMEOBOX PROTEIN {CLON	homeobox protein Gsh-1 [Mus muscul >pir S63604 S63604 homeobox protein mouse >sp P31315 GSHI_MOUSE HC PROTEIN GSH-1. >pir A37290 A372 protein Gsh-1. mouse (fragment) {SU >sp G601819 G601819 HPX HOMEO PROTEIN {CLON	-1 - DBOX meotic 6-205}	gb AAA96814. 1	5157	_	324	65	29	HCRPE63	pSport1
875066 (AF195951) signal recognition particle 68 [Homo sapiens] >sp AAF24308 AAF24308 Signal recognition particle 68. Length = 619	(AF195951) signal recognition particl sapiens] >sp AAF24308 AAF24308 S recognition particle 68. Length = 619	e 68 [Homo ignal	gb AAF24308. 1 AF1959	5158	2	1315	86	66	HCRPE76	pSport1
875067				5159	1	825			HCRPE44	pSport1
875068				5160	2	439			HCRPE34	pSport1
875070 (AF151354) general transcription factor 3 [Homo sapiens] >sp AAF19786 AAF19786 General transcription factor 3. Length = 959	(AF151354) general transcription factor 3 [Fapiens] >sp AAF19786 AAF19786 General transcription factor 3. Length = 959	or 3 [Homo Jeneral	gb AAF19786. 1 AF1513	5161	1	489	96	76	HE8QV20	Uni-ZAP XR

885	875076	(AB007925) KIAA0456 protein [Homo sapiens] >sp 075044 075044 KIAA0456 PROTEIN (FRAGMENT). >gb AAC52480.1  FBP 27 [Mus musculus] {SUB 756-806} Length = 1095	dbj BAA32301  -1	5162	2	577	69	75	нвівQ89	HBIBQ89 Uni-ZAP XR
988	875080	(AB029036) KIAA1113 protein [Homo sapiens] >sp BAA83065 BAA83065 KIAA1113 protein (fragment). Length = 1131	dbj BAA83065 .1	5163	2	958	06	06	HFAAD07	HFAAD07 Uni-ZAP XR
887	875081	DIA-12C protein [Homo sapiens] >emb CAB39108.1  (AL031053) dJ267M20.1 (diaphanous (Drosophila, homolog) 2) [Homo sapiens] {SUB 685-906} Length = 1096	emb CAA7586 9.1	5164	16	576	16	91	H2LAY41	pBluescript SK-
888	875088	(AL031673) dJ694B14.1 (PUTATIVE novel KRAB box protein with 18 C2H2 type Zinc finger domains) [Homo sapiens] >sp Q9Y3M9 Q9Y3M9 DJ694B14.1 (PUTATIVE NOVEL KRAB BOX PROTEIN WITH 18 C2H2 TYPE ZINC FINGER DOMAINS) (FRAGMENT). Length = 738	emb CAB4354 8.1	5165	2	799	98	88	HDPIG12	pCMVSport 3.0
688	875092	(AL132964) putative protein [Arabidopsis thaliana] >sp CAB62464 CAB62464 Hypothetical 189.6 kd protein. Length = 1672	emb CAB6246 4.1	5166	2	385	42	63	HMVCZ67	pSport1
068	875093	male-enhanced antigen-2 [Mus musculus] >splP55937 G160_MOUSE GOLGIN-160 (MALE-ENHANCED ANTIGEN-2) (MEA-2). Length = 1325	dbj BAA19612 .1	5167	2	385	99	59	HWLRF06	pSport1
891	875094	zinc finger protein (mkr5) [Mus musculus] >sp P10078 ZF28_MOUSE_ZINC_FINGER PROTEIN 28 (ZFP-28) (MKR5 PROTEIN) (FRAGMENT). Length = 614	gb AAA37120. 1	5168	2	424	61	99	HTNBJ90	pBluescript SK-
892	875099			5169	133	312			HWLUZ75	pSport1
893	875100	(AK000196) unnamed protein product [Homo sapiens] Length = 239	dbj BAA91003  -1	5170	36	497	26	66	HDTBD43	pCMVSport 2.0
894	875101			5171	-	243			HWLUG07	pSport1

895	875102	[Mus musculus] >pir T30250 T30250 GT1 protein - dbj BAA06184 mouse >sp Q61818 Q61818 HYPOTHETICAL   .1    .1    .1    .1	dbj BAA06184  .1	5172	158	8	69	69	HCRPV30	pSport1
+	875103			5173	6	332			HTPHV54	HTPHV54 Uni-ZAP XR
+	875105			5174	172	426			HWLMY30	pSport1
<del> </del>	875106			5175	29	262			HTTFJ81	Uni-ZAP XR
+	875110			5176	431	826			HDPCC41	pCMVSport 3.0
1	875113			5177	243	473			HINAA28	pSport1
	875114	(AK000388) unnamed protein product [Homo sapiens] Length = 440	dbj BAA91133 .1	5178	3	428	69	74	HTEBS63	Uni-ZAP XR
$\vdash$	875115			5179	384	260			HCROK18	pSport1
	875118		:	5180	3	317			HCROK31	pSport1
	875121	(AK001720) unnamed protein product [Homo sapiens] Length = 605	dbj BAA91860 .1	5181	3	653	98	98	HCROE24	pSport1
	875123	spasmolytic peptide [Homo sapiens] >sp Q03403 SP_HUMAN SPASMOLYTIC POLYPEPTIDE PRECURSOR (SP). Length = 129	gb AAB05397.	5182	80	529	87	87	H2CBN19	pBluescript SK-
<del> </del>	875124			5183	150	329			HDTLM04	pCMVSport 2.0
-	875125			5184	597	445			HOCTE49	pSport1
$\vdash$	875126			5185	129	284			HWLNR78	pSport1
<del> </del>	875131			5186	487	693			НСЕDD96	Uni-ZAP XR
+	875133			5187	3	371			ннғнѕ96	HHFHS96 Uni-ZAP XR
	875134			5188	71	271			HWLN090	pSport1
<del> </del>	875139			5189	79	249			HE2J022	Uni-ZAP XR
-	875143			5190	106	2		1	HCYBJ96	pBluescript SK-

73 HCQDV29 Lambda ZAP	HCRPQ66 pSport1	94 HE9RN07 Uni-ZAP XR	HDQEJ55 pCMVSport 3.0	HCYBJ95 pBluescript SK-	HCUDX92 ZAP Express	HCRON75 pSport1	HWLNR94 pSport1	HCRPY40 pSport1	HHEXW67 pCMVSport	HWLNH10 pSport1	99 HDQEG93 pCMVSport 3.0	88   HWLQT75   pSport1	
63 7		94									66	74 8	,
169	285	979	234	157	218	200	116	846	1299	78	1041	341	
113	100	2	1	2	108	3	3	541	9901	1	133	3	
5191	5192	5193	5194	5195	5196	5197	5198	5199	5200	5201	5202	5203	
emb CAB5182 3.1		emb CAB5307 0.1									gb AAF06998. 1 AF1705	dbj BAA88120 .1	
(AJ222767) ATPase subunit 6 [Cavia porcellus] >sp CAB51823 CAB51823 ATPase subunit 6. Length = 226		(AL035071) dJ1085F17.1.1 (ortholog of mouse DNMT3B (DNA Cytosine-5 Methyltransferase 3B1) [0.1] (isoform 1)) [Homo sapiens] >sb AAD53063.1 AF156488_1 (AF156488) DNA cytosine-5 methyltransferase 3 beta 1 [Homo sapiens] >sp AAD53063 AAD53063 DNA cytosine-5 methyl									(AF170583) ets homologous factor [Homo sapiens] >sp AAF06998 AAF06998 Ets homologous factor. Length = 300	(AB015349) HRHFB2063 [Homo sapiens] >sp BAA88120 BAA88120 HRHFB2063 protein (fragment). Length = 269	
875144	875150	875151	875154	875156	875157	875160	875165	875174	875177	875178	875182	875190	
914	915	916	917	918	919	920	921	922	923	924	925	926	

HCWUO91 ZAP Express	pCMVSport 2.0	Uni-ZAP XR	Uni-ZAP XR	9 pSport1	0 pSport1	5 pSport1	HCQAW68 Lambda ZAP	9 pSport1	Uni-ZAP XR	1 pCMVSport 2.0	2 pCMVSport 3.0	3 pSport1	8 pSport1	3 pBluescript	HMWDC28 Uni-ZAP XR
HCWU09	HDTIP90	HE9TA31	HFPBV89	HWLQZ89	HCRMY90	HNBTB35		HWLRR89	HEICC11	нона из	HHEVA12	HWLPE33	HCRME38	HUSFH63	HMWDC2
100						77	<u>8</u>	99		95		100		84	
100						92	100	57		95		86		83	
179	208	538	464	1407	615	1030	577	318	406	872	344	279	662	582	488
3	77	392	264	1099	304	260	359	961	170	8	165	1	339	166	114
5205	5206	5207	5208	5209	5210	5211	5212	5213	5214	5215	5216	5217	5218	5219	5220
gb AAD16450.						gb AAC83687.	dbjBAA90881 .1	sp P39192 AL US_HUMAN		dbj BAA13508 .1		dbj BAA91324 .1		dbj BAA91263 .1	
(AF102166) intracellular chloride channel CLIC3 [Homo sapiens] >sp O95833 CLI3_HUMAN CHLORIDE INTRACELLULAR CHANNEL PROTEIN 3. Length = 207						(AF083105) HMG box factor SOX-13 [Homo sapiens] >sp 095275 095275 HMG BOX FACTOR SOX-13. Length = 890	(AK000010) unnamed protein product [Homo sapiens] Length = 385	!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!   sp P39192 AL   Length = 585		protein involved in sexual development [Homo sapiens] >sp Q92600 Q92600 CELL DIFFERENTIATION PROTEIN RCD1. Length = 299		(AK000693) unnamed protein product [Homo sapiens] Length = 452		(AK000572) unnamed protein product [Homo sapiens] Length = 328	
875194	875197	875198	875200	875203	875205	875206	875208	875209	875210	875211	875214	875215	875223	875226	875228
828	929	930	931	932	933	934	935	936	937	938	939	940	941	942	943

875236			5221	887	1183			HUVDJ48	Uni-ZAP XR
			5222	209	388			НСОВЕ84	Lambda ZAP II
A & E	(AF023158) tyrosine phosphatase [Homo sapiens] >sp O43183 O43183 TYROSINE PHOSPHATASE (EC 3.1.3.48). Length = 459	gb AAB88293. 1	5223	2	637	84	84	HCYBJ39	pBluescript SK-
			5224	221	364			HCRMW50	pSport1
			5225	612	905			НСQDF84	HCQDF84 Lambda ZAP
F 로	artifact-warning sequence (translated ALU class C) - pir C40201 C4 human Length = 613	pir C40201 C4 0201	5226	241	408	89	74	HNHOD84	HNHOD84 Uni-ZAP XR
			5227	355	636			HACCF57	HACCF57 Uni-ZAP XR
			5228	-	72			HHPGU61	HHPGU61 Uni-ZAP XR
FV 2 9 FT F	put. Cyt repressor (aa 1-341) [Escherichia coli] >gb AAB03066.1  CG Site No. 887 [Escherichia coli] >gb AAC76916.1  (AE000467) regulator for deo operon, udp, cdd, tsx, nupC, and nupG [Escherichia coli] >pir A24963 RPECCT cyt transcription repressor cytr -	emb CAA2731 8.1	5229	72	470	96	96	HFATS83	Uni-ZAP XR
<u>  ∞                                 </u>	similar to G9a gene. [Homo sapiens] >sp Q15047 Q15047 KIAA0067 PROTEIN. Length = 1291	dbj BAA06689 .1	5230	134	916	98	87	HAMFL51	pCMVSport 3.0
			5231	1174	1350			HPLBS64	Uni-ZAP XR
ļ			5232	-	117			HHFGS83	Uni-ZAP XR
			5233	2	214			HCQAI83	Lambda ZAP II
			5234	128	370			HKIAB83	Uni-ZAP XR
2 is	(AK000553) unnamed protein product [Homo sapiens] Length = 298	dbjBAA91249 .1	5235	3	257	86	86	HOUAT80	Uni-ZAP XR

HCUCG82 ZAP Express	pSport1	Lambda ZAP II	pCMVSport 3.0	pSport1	pSport1	pSport1	HCQDD32 Lambda ZAP	pCMVSport 3.0	HCQDT68 Lambda ZAP	Uni-ZAP XR	HAGDP04 Uni-ZAP XR	pSport1	pSport1	pCMVSport 3.0	pCMVSport 3.0	pSport1	pSport1
HCUCG82	HWLMY83	HHGDB82	HHEMA27	HWLQS11	HCRNO87	HCROJ83	нсорр32	HDPQA93	нсорт68	HE2RW42	HAGDP04	HWLRA80	HWLRC80	HWBBH79	HJMAF44	HWLWT47	HWLVG85
													70	64			26
											ļ <u>.</u>	_	64	54			33
345	458	733	207	269	559	438	503	1385	364	853	720	234	364	273	880	695	299
187	279	557	46	09	203	1	192	3	2	617	535	-	125	94	674	408	<b>د</b>
5236	5237	5238	5239	5240	5241	5242	5243	5244	5245	5246	5247	5248	5249	5250	5251	5252	5253
													sp P39194 AL U7_HUMAN	sp P39188 AL  U1_HUMAN			gb AAA91253. 1
													IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593	IIII ALU SUBFAMILY J WARNING ENTRY IIII Length = 591			weak similarity to SP:YAD5_CLOAB (P33746) hypothetical protein and to PIR:C48583 stress-inducible protein ST11 [Caenorhabditis elegans] >pir[T29012 T29012 hypothetical protein ZK328.7 - Caenorhabditis elegans >sp Q23468 Q23468 SIMILARITY TO SP:YAD5_CLOAB.
875279	875280	875281	875282	875287	875288	875292	875296	875303	875304	875305	875306	875307	875308	875309	875310	875311	875312
959	096	961	962	963	964	965	996	296	896	696	970	971	972	973	974	975	976

	а-	~	_		+	t:		<b></b>	<b></b>	4	٠
pSport1	HCQCM79   Lambda ZAP	HMSGP80 Uni-ZAP XR	pSport1	pSportl	HDQFG33 pCMVSport 3.0	pCMVSport 3.0	pSport1	pBluescript SK-	pBluescript SK-	pBluescript	pBluescript SK-
НМУDQ41	НСОСМ79	HMSGP80	HCRNJ78	HWLOY24	НDQFG33	HWBCW80	HCRNL77	H2CBI34	HCYBD76	НКММQ08	HILCJ69
66		42			98						
86		37			84						
433	170	1446	407	258	317	416	9/	292	395	503	381
2	3	145	204	31	မ	246	2	2	15	372	1
5254	5255	5256	5257	5258	5259	5260	5261	5262	5263	5264	5265
emb CAB7077 1.1		emb CAA2011 9.1			dbj BAA22622 .1						
(AL137496) hypothetical protein [Homo sapiens] >emb[CAB70771.1] (AL137496) hypothetical protein [Homo sapiens] >sp[CAB70771]CAB70771 Hypothetical 58.1 kd protein (fragment). Length = 521		(AL031177) dJ889M15.3 (novel protein) [Homo emb CAA2011 sapiens] >sp O95534 O95534 DJ889M15.3 (NOVEL 9.1  PROTEIN) (FRAGMENT). Length = 394			tetracycline transporter-like protein [Mus musculus] >pir JC5641 JC5641 sugar transporter protein HiAT1 - mouse >sp P70187 P70187 HIPPOCAMPUS ABUNDANT PROTEIN TRANSCRIPT 1 (TETRACYCLINE TRANSPORTER-LIKE PROTEIN). Length = 490						
875313	875316	875319	875324	875325	875331	875332	875336	875338	875341	875346	875347
776	978	616	086	981	982	983	984	985	986	286	886

686	875355	regulatory protein [Mus musculus] >emb CAA47648.1  npdcf-1 [Mus musculus] >pir 148691 148691 regulatory protein - mouse >sp Q64322 NPD1_MOUSE NEURAL PROLIFERATION DIFFERENTIATION AND CONTROL PROTEIN-1 PRECURSOR (NPDC-1 PROTEIN). Length = 332	gb AAA39836. 1	5266	464	1138	78	83	HDPGF81	pCMVSport 3.0
066	875356	regulatory protein [Mus musculus] >emb CAA47648.1  npdcf-1 [Mus musculus] >pir 148691 148691 regulatory protein - mouse >sp Q64322 NPD1_MOUSE NEURAL PROLIFERATION DIFFERENTIATION AND CONTROL PROTEIN-1 PRECURSOR (NPDC-1 PROTEIN). Length = 332	gb AAA39836. 1	5267	30	341	68	91	HUSGQ41	pSport1
166	875360			5268	711	896			HPMFC89	Uni-ZAP XR
992	875364			5269	429	662			HWLWK37	pSport1
993	875366	(AF050078) growth arrest specific 11 [Homo sapiens] >gb AAC69519.1  (AF050079) growth arrest specific 11 [Homo sapiens] >sp O95995 O95995 GROWTH ARREST SPECIFIC 11. Length = 478	gb AAC69518. 1	5270	74	211	95	26	HSYAG49	pCMVSport 3.0
994	875367	ALU SUBFAMILY J WARNING ENTRY       Length = 591	sp P39188 AL U1_HUMAN	5271	152	298	62	9	HAGFQ75	HAGFQ75 Uni-ZAP XR
995	875371	(AK001527) unnamed protein product [Homo sapiens] Length = 790	dbj BAA91741 .1	5272	94	297	98	98	нснм074	pSport1
966	875372			5273	32	220			HCQCL42	Lambda ZAP II
766	875373	apomucin [Homo sapiens] >pir A57534 A57534 mucin (clone L31) - human (fragment) >sp Q13792 Q13792 APOMUCIN (FRAGMENT). Length = 1042	emb CAA8830 7.1	5274	2	772	97	76	HHFOB15	Uni-ZAP XR
866	875377			5275	94	267			HCRMB64	pSport1
666	875378	[AJ245539] GalNAc-T5 [Homo sapiens] >sp CAB65104 CAB65104 GalNAc-T5 (fragment). Length = 668	emb CAB6510 4.1	5276	1	615	66	66	H2LAB72	pBluescript SK-

HE80D44 Uni-ZAP XR	pSport1	pSport1	pSport1	Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1	HFXKG78 Lambda ZAP	Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	HCQCX73 Lambda ZAP	pSport1	Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	Lambda ZAP II	HCQDN71 Lambda ZAP
HE80D44	HCRMZ16	HWLMZ75	HWLMT21	HCEMB73	HWLNF24	HNHNC74	HCRNF23	HFXKG78	HFPFG11	HCROG59	HLYBH74	HBGNK79	нсосх73	ELDOTMH	HMSIB72	HWLMC85	HCRNH72	HSDHD72	HCQAB70	HCQDN71
50								82				88								
27								82				88								
687	432	365	503	959	655	183	544	1790	358	581	220	969	328	230	190	328	991	462	238	386
124	274	189	252	315	485	-	569	120	230	153	77	1	53	111	41	158	872	316	2	210
5277	5278	5279	5280	5281	5282	5283	5284	5285	5286	5287	5288	5289	5290	5291	5292	5293	5294	5295	5296	5297
gb AAD56247.  1 AF1864								dbj BAA13437 .1				emb CAB5305 8.1								
(AF186461) ring finger protein Fxy [Rattus norvegicus] >sp AAD56247 AAD56247 Ring finger protein Fxy. Length = 667								KIAA0242 protein [Homo sapiens] >sp Q92575 Q92575 MYELOBLAST KIAA0242 (FRAGMENT). Length = 529				(AL031033) C321D2.4 (novel protein) [Homo sapiens] >splCAB53058 CAB53058 C321D2.4 (novel protein) (fragment). Length = 262								
875379	875380	875381	875382	875384	875385	875388	875391	875397	875402	875405	875406	875410	875415	875416	875417	875418	875419	875423	875425	875427
1000	1001	1002	1003	1004	1005	1006	1007	1008	1009	1010	1011	1012	1013	1014	1015	1016	1017	1018	1019	1020

Lambda ZAP II	HCQAW10 Lambda ZAP	pSport1	pSport1	Uni-ZAP XR	pBluescript SK-	Uni-ZAP XR	pSport1	pSport1	pCMVSport 2.0	HTHDF09 Uni-ZAP XR	pCMVSport 2.0	pSport1	pSport1	pSport1
HCQCQ73 Lambda ZAP	HCQAW10	HCRNE71	HWLNY71	HTXSH02	H2CBL70	HNFFQ01	HCRMD70	HWLWX54	HDTBL01	нтнрғ09	НОНАD26	HWLQB70	HCRNJ70	HCHAN69
86					83	95			87		66		100	
97					79	95			87		66		100	
550	454	280	822	396	816	184	290	330	558	231	2112	500	358	520
2	251	7	865	151	_	80	294	133	115	1	_	3	11	221
5298	5299	5300	5301	5302	5303	5304	5305	5306	5307	5308	5309	5310	5311	5312
gb AAD51455.					dbj BAA11492 .1	gb AAC39892. 1			emb CAB7076 8.1		gb AAC12926.		gb AAD45398. 1	
(AC004955) supported by ESTs T61992 (NID:g665235) and W26450 (NID:g1307167) and Genscan [Homo sapiens] >sp AAD51455 AAD51455 WUGSC:H_DJ1087M19.1 protein. Length = 557					similar to protein kinase of X.laevis, has putative transmembrane domain incentral region [Homo sapiens] >sp Q14680 Q14680 KIAA0175 PROTEIN. Length = 651	(AF047441) RNA polymerase I 40kD subunit [Homo gb AAC39892. sapiens] >sp O75395 O75395 RNA POLYMERASE 1  140kD SUBUNIT. Length = 342			(AL137489) hypothetical protein [Homo sapiens] >emb CAB70768.1  (AL137489) hypothetical protein [Homo sapiens] >sp CAB70768 CAB70768 Hypothetical 12.7 kd protein (fragment). Length = 116		(AF037448) Gry-rbp [Homo sapiens] >sp O60506 O60506 GRY-RBP. Length = 623		(AF086709) NAG-7 protein [Homo sapiens] >sp Q9Y6C7 Q9Y6C7 NAG-7 PROTEIN. Length = 94	
875428	875429	875433	875434	875437	875440	875441	875442	875446	875452	875458	875460	875461	875462	875463
1021	1022	1023	1024	1025	1026	1027	1028	1029	1030	1031	1032	1033	1034	1035

pCMVSport 3.0	pBluescript SK-	pSport1	pSport1	pSport1	pSport1	Uni-ZAP XR	Uni-ZAP XR	Jni-ZAP XR	Jni-ZAP XR	pCMVSport 2.0	pSport1	Lambda ZAP II	pCMVSport 3.0	pBluescript SK-	pSport1	pSport1	pSport1	pSport1
HDPXJ69	H2CBP05	910NTMH	HCROC40	HWLWW31	HWLOU12	HPTTL69	HT3BA65	HMSHD68 Uni-ZAP XR	HSUAE53 Uni-ZAP XR	HTJMN69	HHMMD68	HCQDM23 Lambda ZAP	ннемо68	H2CBM67	HWLWJ34	HWLRL54	HCROI48	HCRMM67
												7 97						
406	872	354	498	377	197	396	273	260	716	265	296	534 97	292	609	344	387	268	524
218	99	1	250	147	3	-	130	72	456	29	3	-	140	460	21	187	74	336
5313	5314	5315	5316	5317	5318	5319	5320	5321	5322	5323	5324	5325	5326	5327	5328	5329	5330	5331
												pir A49963 A4 3932						
												mucin 2 precursor, intestinal - human (fragments) pir A49963 A4						
875468	875474	875475	875477	875478	875479	875481	875484	875486	875490	875491	875492	875493	875495	875496	875498	875499	875500	875501
1036	1037	1038	1039	1040	1041	1042	1043	1044	1045	1046	1047	1048	1049	1050	1051	1052	1053	1054

1055	875502	NEDD1 PROTEIN (FRAGMENT). >dbj BAA01554.1   nedd-1 protein [Mus musculus] {SUB 16-675} Length = 675	sp P33215 NE D1_MOUSE	5332	2	1213	79	88	HTFNZ86	pSport1
1056	875503			5333	205	402			HCNCD90	HCNCD90 Lambda ZAP
1057	875508	(AB002334) KIAA0336 [Homo sapiens] >sp O15045 O15045 KIAA0336. Length = 1583	dbj BAA20794 .1	5334	2	775	87	68	HMVDK54	pSport1
1058	875512			5335	138	317			нсосу65	HCQCV65 Lambda ZAP
1059	875514			5336	2	373			99ANTMH	pSport1
1060	875515	(AB023201) KIAA0984 protein [Homo sapiens] >sp Q9Y219 Q9Y219 KIAA0984 PROTEIN (FRAGMENT). Length = 728	dbj BAA76828 .1	5337	17	424	100	100	HLYC165	pSport1
1061	875516			5338	2	361			HKAA067	pCMVSport 2.0
1062	875517			5339	126	260			HCE3W64	Uni-ZAP XR
1063	875518	LIKE	sp P17082 RR A2_HUMAN	5340	22	885	95	95	HKAKX87	pCMVSport 2.0
1064	875520	GATA-6 [Homo sapiens] >gb AAC50941.1  hGATA-6 [Homo sapiens] >sp Q92908 GAT6_HUMAN TRANSCRIPTION FACTOR GATA-6 (GATA BINDING FACTOR-6). Length = 449	dbj BAA22621 .1	5341		837	66	66	HUSGX12	pSport1
1065	875523			5342	1	165			HCNDZ15	HCNDZ15 Lambda ZAP
1066	875525	catalase [Campylobacter jejuni] >pir 140767 140767 catalase (EC 1.11.1.6) - Campylobacter jejuni >sp Q59296 CATA_CAMJE CATALASE (EC 1.11.1.6). Length = 507	emb CAA5944	5343	306	130	84	68	HCFNM40	pSport1
1067	875527	(AK001870) unnamed protein product [Homo sapiens] Length = 278	dbj BAA91953 .1	5344	261	569	93	94	HMSGC65	HMSGC65 Uni-ZAP XR

HCQDN81 Lambda ZAP	pSport1	pSport1	pSport1	pSport1	pSport1	pCMVSport 1	pCMVSport 3.0	HCQAF61 Lambda ZAP	HCQCX63 Lambda ZAP	pSport1	Uni-ZAP XR	pBluescript SK-	pSport1
HCQDN81	HFICY86	HNTSA70	HWLMX64	HTWFG63	HWLNY32	HLJDL64	ннеом62	HCQAF61	нсосх63	HOVET54	HRODW53	H2CBE60	HWMCK45
		96				74	06				100	66	
		96				71	06				66	86	
155	9/9	875	100	162	241	333	555	420	207	059	455	597	318
3	389	E.	2	-	2	175		193	22	411	108	46	88
5345	5346	5347	5348	5349	5350	5351	5352	5353	5354	5355	5356	5357	5358
		dbj BAA12699 .1				emb CAA2403 1.1	gb AAC39837.   1		·		dbj BAA91139 .1	dbj BAA11502 .1	
		HsMcm6 [Homo sapiens] >sp Q14566 MCM6_HUMAN DNA. REPLICATION LICENSING FACTOR MCM6 (P105MCM). Length = 821				ATPase 6 [Homo sapiens] >gb AAB58948.1  ATPase 6 [Homo sapiens] >pir A01049 PWHU6 H+- 11.1  transporting ATP synthase (EC 3.6.1.34) protein 6- human mitochondrion >sp P00846 ATP6_HUMAN ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6). Length = 226	(AF059531) protein arginine N-methyltransferase 3 [Homo sapiens] >sp O60678 O60678 PROTEIN ARGININE N-METHYLTRANSFERASE 3 (FRAGMENT). Length = 512				(AK000399) unnamed protein product [Homo sapiens] Length = 427	similar to hypothetical protein YM9959.11C of S.cerevisiae. [Homo sapiens] >sp[Q14690 RRP5_HUMAN RRP5 PROTEIN HOMOLOG (KIAA0185) (FRAGMENT). Length = 1884	
875528	875529	875534	875538	875539	875543	875544	875545	875546	875547	875548	875550	875551	875552
1068	1069	1070	1071	1072	1073	1074	1075	1076	1077	1078	6201	1080	1081

				T	T-					_				
ď	pSport1	HTLEY14 Uni-ZAP XR	pCMVSport 2.0	Uni-ZAP XR	HCQAG54 Lambda ZAP	pSport1	pSport1	pSport1	pBluescript SK-	pSport1	pSport1	pCMVSport 3.0	pSport1	pSport1
HKAFL60	HUSXP66	HTLEY14	HOFMV44	HSLJN60	HCQAG54	HHMMD60	HWLMB59	HUFAU68	H2LAX58	HCRQD82	HCRPV05	HHECM62	HFOXW88	HWLXT17
		86	58	88				100	82					62
		86	39	84				100	08					50
262	438	627	1138	392	365	441	1038	1081	130	151	323	330	542	715
14	193	328	2	8	150	256	662	2	2	524	183	148	375	2
5359	5360	5361	5362	5363	5364	5365	9988	5367	5368	5369	5370	5371	5372	5373
		gb AAA67217. 1	dbj BAA92121 .1	gb AAB70111. 1				dbj BAA86499 .1	\$6Y5 \$6Y5					gb AAF24125. 1 AF1217
		a peptide [Cloning vector	(AK002174) unnamed protein product [Homo sapiens] Length = 508	(AC002542) similar to C. elegans F11A10.5; 80% similarity to Z68297 (PID:g1130619) [Homo sapiens] >sp O14577 O14577 WUGSC:H_RG114A06.1 PROTEIN. Length = 434				S	IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414					(AF121775) nasopharyngeal carcinoma susceptibility gb AAF24125. protein LZ16 [Homo sapiens]
875553	875554	875556	875558	875559	875560	875563	875564	875565	875567	875570	875572	875573	875574	875578
1082	1083	1084	1085	1086	1087	1088	1089	1090	1091	1092	1093	1094	1095	1096

HODAY72 Uni-ZAP XR	HCQBIS6 Lambda ZAP	HTTCM45 Uni-ZAP XR	HARNM58 pCMVSport 3.0	HMIAQ09 Uni-ZAP XR	HE9MD57 Uni-ZAP XR	HCQDA63 Lambda ZAP	pSport1	pCMVSport 3.0	HMUBG89 pCMVSport 3.0	pCMVSport 3.0	pSport1	HROBR56 Uni-ZAP XR	pSport1	pSport1	pSport1	Uni-ZAP XR
HODAY72	нсові56	HTTCM45	HARNM58	HMIAQ09	HE9MD57	нсоруе н	HWLR057	инеооео	HMUBG89	HDPRN70	HCRMC33	HROBR56	HWLMU33	HCRQC94	HCRMQ55	HSAZF81
														2 92		
299	164	438	612	1019	332	1150	564	452	692	355	446	351	402	660 92	624	695
434 (	س ا	283 4	367	150 1	156	695 1	337	303	371	134	234 4	130	211 4	1	430 (	207
5374	5375	5376	5377	5378	5379	5380	5381	5382	5383	5384	5385	5386	5387	5388	5389	5390
														dbj BAA09764 .1		
														The KIAA0143 gene product is related to a putative of C.elegans gene encoded on cosmid C32D5. [Homo sapiens] >sp Q14156 Y143_HUMAN HYPOTHETICAL PROTEIN KIAA0143 (FRAGMENT). Length = 885		
875583	875584	875585	875587	875588	875589	875590	875594	875596	875597	875598	875600	875604	875605	875606	875608	875609
1097	1098	1099	1100	1101	1102	1103	1104	1105	1106	1107	1108	1109	1110		1112	1113

HTJMO37 pCMVSport 2.0	pBluescript	pSport1	pCMVSport 3.0	pBluescript SK-	HCQCX54 Lambda ZAP	HCQCG75 Lambda ZAP	pCMVSport 3.0	Other	pSport1	pCMVSport 3.0	HKMAB82 Uni-ZAP XR	Uni-ZAP XR
HTJM037	HKCSA54	HWLQA55	HWBDT63	Н2СВQ54	нсосх54	нсосс75	HHEZN36	HPCIS18	HISAT54	HLWAC54	HKMAB82	HPVAB96
			99					68			95	<i>L</i> 9
			43					68			98	50
397	430	206	653	657	252	377	630	189	009	272	145	302
251	161	09	267	26	28	135	382	_	436	3	2	က
5391	5392	5393	5394	5395	5396	5397	5398	5399	5400	5401	5402	5403
			gb AAA88038. 1					gb AAC50379.			gb AAC40188. 1	dbj BAA83026 .1
			protein [Homo sapiens] >sp Q14288 Q14288 HYPOTHETICAL PROTEIN (FRAGMENT). Length = 641					TAR RNA loop binding protein [Homo sapiens] >pir S62356 S62356 TRP-185 protein - human >sp Q13395 Q13395 TAR RNA LOOP BINDING PROTEIN. Length = 1621			CID 2 4 5 (4 5 )	(AB028997) KIAA1074 protein [Homo sapiens] >sp BAA83026 BAA83026 KIAA1074 protein. >dbj BAA91516.1  (AK001137) unnamed protein product [Homo sapiens] {SUB 1-546} >emb CAB70706.1  (AL137351) hypothetical protein [Homo sapiens] {SUB 1337-1709} Length = 1709
875610	875611	875612	875613	875625	875628	875629	875630	875631	875632	875633	875634	875635
1114	1115	1116	1117	1118	1119	1120	1121	1122	1123	1124	1125	1126

					10			· · · · · · · · · · · · · · · · · · ·		<u>-</u>
HBMSX53 Uni-ZAP XR	pSport1	Uni-ZAP XR	HMWFZ60 Uni-ZAP XR	pSport1	ZAP Express	pSport1	pSport1	pSport1	pSport1	HOEEYS3 Uni-ZAP XR
HBMSX53	HCFCS58	HPMK129	HMWFZ60	HUCPH16	HCUDA52	HTWCN56	HWLUF58	HWLMI53	HWLMB54	НОЕЕҮ53
		68	93	91			82	78		
		87	93	91			82	92		
226	2228	551	905	896	288	81	669	481	122	655
113	2043	E.	639	6	76	1	85	∞	3	449
5404	5405	5406	5407	5408	5409	5410	5411	5412	5413	5414
		emb CAB3798 1.1	.1	gb AAC96102. 1			dbj BAA87339 .1	dbj BAA87339 .1		
		(AL022395) dJ273N12.1 (PUTATIVE protein based on EST matches) [Homo sapiens]       1.1          >gb AAF04511.1 AF174590_1 (AF174590) F-box protein Fbl4 [Homo sapiens] >sp O95919 O95919       .         DJ273N12.1 (PUTATIVE PROTEIN BASED ON EST MATCHES) (FRAGMENT).       >sp AAF04511 AAF04511 F-bo	(AB032253) bromodomain adjacent to zinc finger domain 1B [Homo sapiens] >dbj BAA89210.1  (AB032253) bromodomain adjacent to zinc finger domain 1B [Homo sapiens] >sp BAA89210 BAA89210 Bromodomain adjacent to zinc finger domain 1B. Length = 1527	(AF071771) SPH-binding factor [Homo sapiens] Length = 551			(AB023416) ASC [Homo sapiens] >sp BAA87339 BAA87339 ASC protein. Length = 195	(AB023416) ASC [Homo sapiens] >sp BAA87339 BAA87339 ASC protein. Length = 195		
875636	875638	875639	875640	875641	875642	875646	875650	875651	875653	875654
1127	1128	1129	1130	1131	1132	1133	1134	1135	1136	1137

pSport1	pSport1	pSport1	Uni-ZAP XR	pSport1	pSport1	Lambda ZAP II	pSport1	pSport1	Uni-ZAP XR	pSport1	pBluescript SK-	Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR
нисос25	HCRMS71	HWLMS13	HE6GF82	HSPBC14	HOCNE41	нсовея	HWLMX40	HCRMB51	нсввн61	HCRNZ51	H2CAA51	HT3AI55	HLWBA37	НЕ2ГР33
66								68		73		86		
86								88		63		86		
558	955	341	554	108	419	266	197	265	533	98	540	504	244	332
-	440	180	282	-	132	159	3	99	246	12	295	1	2	165
5415	5416	5417	5418	5419	5420	5421	5422	5423	5424	5425	5426	5427	5428	5429
emb CAB7073								dbj BAA90899  .1		gb AAB50206. 1		dbj BAA91018 .1		
(AL137442) hypothetical protein [Homo sapiens] >emb CAB70739.1  (AL137442) hypothetical protein [Homo sapiens] >sp CAB70739 CAB70739 Hypothetical 34.5 kd protein (fragment). Length = 316								(AK000040) unnamed protein product [Homo sapiens] Length = 387		[Homo sapiens] >sp[Q99770 Q99770 HYPOTHETICAL 15.4 KD PROTEIN. Length = 139		(AK000219) unnamed protein product [Homo sapiens] Length = 420		
875658	875661	875662	875663	875665	875669	875672	875673	875677	875678	875680	875681	875682	875683	875687
1138	1139	1140	1141	1142	1143	1144	1145	1146	1147	1148	1149	1150	1151	1152

pSport1	pBluescript	pSport1	Lambda ZAP II	pSport1	HRGDD40 Uni-ZAP XR	pBluescript SK-	HMSGN49 Uni-ZAP XR	pSport1	Other	Lambda ZAP II
HCRMN10	HKMMR61	HUFDC50	HKLAB51	HCGBB63	HRGDD40	H2LAD49	HMSGN49	HWLMC49	HAVME52	<b>НС</b> QD <b>Р</b> 49
91				100	50	93		62		28
91				100	41	93		79		44
401	1105	347	280	619	965	154	188	490	1390	511
216	707	156	4	71	402	11	3	2	1187	2
5430	5431	5432	5433	5434	5435	5436	5437	5438	5439	5440
dbj BAA74430 .1				gb AAD42056. 1 AF0449	dbj BAA28677 .1	sp Q9Y6Y5 Q9 Y6Y5		dbj BAA91388 .1		gb AAD21038. 1
(AB021638) X11-like protein 2 [Homo sapiens] >gb AAC72275.1  (AC005954) mint 3 [Homo sapiens]; X11gamma [Homo sapiens] >pir JG0181 JG0181 X11L2 protein - human >sp O96018 APB3_HUMAN AMYLOID BETA A4 PRECURSOR PROTEIN-BINDING FAMILY A MEMBER 3 (NEURON- SPEC				(AF044953) NADH:ubiquinone oxidoreductase PGIV subunit [Homo sapiens] >sp Q9Y6N0 Q9Y6N0 NADH:UBIQUINONE OXIDOREDUCTASE PGIV SUBUNIT. Length = 172	(AB015041) PIF1 [Caenorhabditis elegans] >pir[T37310]T37310 PIF1 protein - Caenorhabditis elegans >sp O61299 O61299 PIF1. Length = 677	IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414		(AK000820) unnamed protein product [Homo sapiens] Length = 122		(AF113131) host cell factor homolog LCP [Homo sapiens] >dbj BAA91898.1  (AK001771) unnamed protein product [Homo sapiens] >sp Q9Y2U9 Q9Y2U9 HOST CELL FACTOR HOMOLOG LCP. Length = 406
875688	875689	875690	875697	875698	875699	875700	875703	875704	875705	875708
1153	1154	1155	1156	1157	1158	1159	1160	1161	1162	1163

pSport1	pCMVSport 3.0	pSport1	Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	HODBC46 Uni-ZAP XR	pBluescript SK-	ZAP Express	pSport1	pSport1	pSport1	pBluescript SK-	Lambda ZAP II	pCMVSport 2.0	HCQDI44   Lambda ZAP II
HCROW44	HDPHF03	HCRM082	HFCDF47	HFICJ16	HWLLU74	HLMDL53	HODBC46	HCYB046	HCUEB32	HCRNQ45	HWL0086	HSPME53	H2CBE48	нсорл47	HDTKC01	нсорі44
81								86					100			
81								93					86			
575	265	627	1133	869	482	823	174	485	808	260	726	387	267	318	138	194
216	101	553	273	528	183	470	10	192	251	141	520	1	16	145	1	3
5441	5442	5443	5444	5445	5446	5447	5448	5449	5450	5451	5452	5453	5454	5455	5456	5457
gb AAC33006. 1								gb AAC79987. 1					emb CAA0646 2.1			
(AF074264) LDL receptor-related protein 6 [Homo sapiens] >pir JE0272 JE0272 low density lipoprotein receptor-related protein 6 - human >sp O75581 O75581 LDL RECEPTOR-RELATED PROTEIN 6. Length = 1613								(AF060219) RCC1-like G exchanging factor RLG [Homo sapiens] >sp 095199 095199 RCC1-LIKE G EXCHANGING FACTOR RLG. Length = 551					(AJ005273) Kin17 [Homo sapiens] >sp O60870 O60870 KIN17 PROTEIN. Length = 393			
875717	875719	875722	875724	875725	875727	875728	875729	875731	875733	875734	875736	875737	875738	875739	875740	875746
1164	1165	1166	1167	1168	1169	1170	1171	1172	1173	1174	1175	1176	1177	1178	1179	1180

HNFGP44 Uni-ZAP XR	pSport1	pSport1	HCQAC43   Lambda ZAP II	pSport1	pSport1	pBluescript SK-	HWTCF43   Uni-ZAP XR	pSport1	HCQDD42 Lambda ZAP	pSport1	pSport1	pCMVSport 3.0	Uni-ZAP XR	HMCIK65 Uni-ZAP XR	HDTGQ43 pCMVSport 2.0
HNFGP44	HWLQG44	HHMMD44	HCQAC43	HWLUF33	HCRPE66	HCYBD73	HWTCF43	HCRNA26	нсорр42	HCRNN21	HCRNH26	HDP:WD42	HTAET42	HMCIK65	HDTGQ43
9/							09								
75							49								
330	280	207	283	450	200	441	2	366	488	238	812	124	243	808	316
112	2	58	83	265	321	310	61	193	240	2	78	38	1	059	2
5458	5459	5460	5461	5462	5463	5464	5465	5466	5467	5468	5469	5470	5471	5472	5473
emb CAB6611 8.1							gb AAB02291. 1								
(AL050348) dJ447F3.2 (ubiquitin-conjugating enzyme E2 H10) [Homo sapiens] >gb AAB53362.1  cyclin-selective ubiquitin carrier protein [Homo sapiens] >sp O00762 UBCB_HUMAN UBIQUITIN-CONJUGATING ENZYME E2 H10 (EC 6.3.2.19) (UBIQUITIN-PROTEIN LIGASE) (UBIQUIT							reverse transcriptase [Homo sapiens] Length = 361								
875747	875751	875752	875753	875754	875760	875761	875765	875766	875768	875769	875772	875773	875774	875778	875779
1181	1182	1183	1184	1185	1186	1187	1188	1189	1190	1191	1192	1193	1194	1195	1196

875780	GD3 synthase [Homo sapiens] >gb AAC37586.1  ganglioside-specific alpha-2, 8-polysialyltransferase [Homo sapiens] >pir A54032 A54032 alpha-N-acetylneuraminate alpha-2,8-sialyltransferase (EC 2.4.99.8) - human >sp Q92185 CAG8_HUMAN ALPHA-N-ACETYL-NEURAMINNI	emb CAA5489	5474	498	863	87	88	HT2SF78	Uni-ZAP XR
	ZK520.1 [Caenorhabditis elegans] >pir[T27880 T27880 hypothetical protein ZK520.1 - Caenorhabditis elegans >sp O46018 O46018 ZK520.1 PROTEIN. Length = 519	emb CAB0729 9.1	5475	2	712	38	61	HCRMG60	pSport1
875782			5476	19	492			HCRNC13	pSport1
875783			5477	142	264			HCRPH74	pSport1
875784			5478	92	283			HCQDW41	Lambda ZAP II
875785			5479	175	318			HCRMZ22	pSport1
875786			5480	220	423			нсоре41	Lambda ZAP II
875787			5481	230	532			HMKCZ06	pSport1
875789			5482	806	1168			HMEGG05	HMEGG05 Lambda ZAP
875792			5483	156	332			HNTMD41	pSport1
875794	(AK002156) unnamed protein product [Homo sapiens] Length = 326	dbj BAA92113 .1	5484	2	721	\$6	97	HCRNJ24	pSport1
875798			5485	315	455			HWABK33	pCMVSport 3.0
875800			5486	282	497			HCYBC44	pBluescript SK-
875801			5487	283	507			HWLQA40	pSport1
875804			5488	1	330			HWHPI43	pCMVSport 3.0
875805			5489	25	390			HKCSF43	pBluescript

1213	875808		KIC A A 0/404	5490	462	752	90		š	}
1214	875809	(AJ001714) Myosin-IXA [Homo sapiens] emt >sp CAA04947 CAA04947 Myosin-IXA (fragment). 7.1  Length = 774	emb CAA0494 7.1	5491	20.7	1088	3	દ		HCKNL08
1215	875810	•		5492	2	256			H	HCRNY14
1216	875814	(AF105376) heparan sulfate D-glucosaminyl 3-O-sulfotransferase-3A [Homo sapiens] >splQ9Y663 Q9Y663 HEPARAN SULFATE D-GLUCOSAMINYL 3-O-SULFOTRANSFERASE-3A (EC 2.8.2.23). Length = 406	gb AAD30208. 1 AF1053	5493	8	824	52	54	H	HCRQG46
1217	875815			5494	226	423			H	HCRQK63
1218	875816			5495	149	466			HМ	HWLVS38
1219	875817	(AL035461) d1967N21.5 (novel MCM2/3/5 family member) [Homo sapiens] >splCAB55276 CAB55276   D1967N21.5 (novel MCM2/3/5 family member) (fragment). Length = 606	emb CAB5527 6.1	5496	44	556	16	94	HC	HCRNT27
1220	875819			5497	12	83			HCR	HCRMT24
1221	875820			5498	1	222			HCR	HCRNQ33
1222	875821			5499	22	480			HMI	HWLU071
1223	875822	ARG5,6 [Candida albicans] >sp P78586 AR56_CANAL_ARG5,6 PROTEIN PRECURSOR [CONTAINS: N-ACETYL-GAMMA-GLUTAMYL-PHOSPHATE REDUCTASE (EC 1.2.1.38) (N-ACETYL-GLUTAMATE SEMIALDEHYDE DEHYDROGENASE) (NAGSA DEHYDROGENASE); ACETYLGLUTAMATE KINASE (EC 2.7.2.8) (NAG K	emb CAA6738 3.1	5500	336	761	31	45	HTX	HTXRZ02
1224	875824			5501	3	302			HWI	HWMB047
1225	875825	(AF156551) putative E1-E2 ATPase [Mus musculus] gb AAF09449.   sp AAF09449 AAF09449 Putative E1-E2 ATPase.   I AF1565   Length = 1187	gb AAF09449. 1 AF1565	5502	3	497	63	78	ЭЭН	нсосс37

HUVGY13 Uni-ZAP XR	HPMFMS9 Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1	pCMVSport 3.0	HTGBQ40 Uni-ZAP XR	pCMVSport 3.0	pSport1	pCMVSport 3.0	pSport1	pSport1	pSport1	Uni-ZAP XR	HCQCL72 Lambda ZAP	HCQCT09 Lambda ZAP	pSport1	pSport1
HUVGY13	HPMFMS9	HCROI42	HACBB04	HMMAC34	HDPFA20	HTGBQ40	HDPWD53	HCROZ63	HWABJ67	HCRMY91	HNTRA39	HCRPW33	HFCF137	HCQCL72	нсост09	HCRMR12	HCIAE18
	74		99														
		_	65														
450	662	458	930	410	522	162	329	360	547	865	623	565	187	214	1016	123	564
289	8	126	460	276	49	-	177	1	347	572	441	437	68	11	732	1	100
5503	5504	5205	5506	2507	5508	5509	5510	5511	5512	5513	5514	5515	5516	5517	5518	5519	5520
	emb CAB5367 7.1		gb AAF05761. 1 AF1925														
	(AL110217) hypothetical protein [Homo sapiens] >emb CAB53677.1  (AL110217) hypothetical protein [Homo sapiens] >pir[T14757[T14757 hypothetical protein DKFZp572C163.1 - human (fragment) >sp CAB53677 CAB53677 Hypothetical 80.6 kd protein (fragment). Length		(AF192529) RPA-binding trans-activator [Homo sapiens] >sp AAF05761 AAF05761 RPA-binding trans-activator. Length = 196														
875826	875828	875832	875833	875834	875836	875837	875838	875839	875840	875841	875845	875846	875848	875849	875850	875851	875852
1226	1227	1228	1229	1230	1231	1232	1233	1234	1235	1236	1237	1238	1239	1240	1241	1242	1243

HHFHU39 Uni-ZAP XR	HCQAW29 Lambda ZAP	pBluescript	pBluescript	pCMVSport	pSport1	pBluescript SK-	pSport1	Lambda ZAP II	pCMVSport 3.0	pSport1	HNHHM31 Uni-ZAP XR	pSport1
ннғн 039	HCQAW29	HBMDM33	HKLSD32	HYACE34	HNTTC18	H2CAA34	HWLQA33	нсост65	HWHPI50	HCRQD12	HNHHM31	HCRQG23
				50	100	84						100
				33	86	82						100
415	264	605	202	516	230	1981	410	263	2653	408	460	341
137	121	3	2	-	E.	788	177	24	2189	220	56	က
5521	5522	5523	5524	5525	5526	5527	5528	5529	5530	5531	5532	5533
				emb CAB0654 5.1	gb AAA79359. 1	gb AAF09482. 1 AF1910						emb CAA4774 9.1
				predicted using Genefinder; cDNA EST yk469a11.5 comes from this gene [Caenorhabditis elegans] >emb CAB01706.1  predicted using Genefinder; cDNA EST yk469a11.5 comes from this gene [Caenorhabditis elegans] >pir T21387 T21387 hypothetical protein F26A3.7 -	DNA binding protein [Homo sapiens] >sp P51523 ZN84_HUMAN ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2). >pir B32891 B32891 finger protein 2, placental - human {SUB 88-738} >sp G238102 G238102 ZINC FINGER. {SUB 71-257} Length = 738	(AF191018) E2IG3 [Homo sapiens] >sp AAF09482 AAF09482 E2IG3. Length = 560						polypeptide BM28 [Homo sapiens] >pir S42228 S42228 replication licensing factor MCM2 - human Length = 892
875855	875856	875858	875863	875864	875865	875868	875871	875874	875884	875886	875888	875891
1244	1245	1246	1247	1248	1249	1250	1251	1252	1253	1254	1255	1256

pBluescript	pBluescript SK-	Lambda ZAP II	pBluescript SK-	pSport1	pCMVSport 3.0	HCQAH30 Lambda ZAP	pCMVSport 3.0	HCQAM30 Lambda ZAP	HAGEA31 Uni-ZAP XR	pSport1	pCMVSport 3.0	pCMVSport 2.0	Uni-ZAP XR
HKLSB39	H2CBN05	нсорт85	HARAJ31	HCRMQ35	HMUBG30	<b>НС</b> QАН30	HWDAH30 pCMVSport 3.0	<b>НС</b> QАМ30	HAGEA31	HCROZ66	HDPBY50	HDTKD18	HHPGT16
79	99						100		92				83
29	47						93		63				83
540	846	224	192	554	341	181	189	252	156	845	829	1353	623
1	355	09	-	351	3	83	-	<i>L</i> 9	-	618	479	1153	1567
5534	5535	5536	5537	5538	5539	5540	5541	5542	5543	5544	5545	5546	5547
pir A32891 A3 2891	pir JC4296 JC4 296						emb CAB6507		gb AAD34084. 1 AF1518				dbj BAA20828 .1
finger protein 1, placental - human >sp P51522 ZN83_HUMAN ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1). Length = 428	ring finger protein - fruit fly (Drosophila melanogaster) Length = 222						(AJ131890) DNA polymerase lambda [Homo sapiens] >gb AAF27541.1 AF161019_1 (AF161019) 4.1  DNA polymerase beta-N [Homo sapiens] >sp CAB65074 CAB65074 DNA polymerase lambda. >sp AAF27541 AAF27541 DNA polymerase beta-N. Length = 575		(AF151847) CGI-89 protein [Homo sapiens] >sp Q9Y397 Q9Y397 CGI-89 PROTEIN. Length = 382				(AB002371) KIAA0373 [Homo sapiens] >sp O15078 O15078 KIAA0373. Length = 1539
875894	875897	875899	875900	875904	875905	875906	875907	875908	875912	875913	875914	875915	875923
1257	1258	1259	1260	1261	1262	1263	1264	1265	1266	1267	1268	1269	1270

pBluescript SK-	Lambda ZAP II	HUKFO71 Lambda ZAP	HCQAT28 Lambda ZAP	pBluescript SK-	pSport1	Uni-ZAP XR	pCMVSport 3.0	pSport1	HSIDV66 Uni-ZAP XR	pCMVSport 3.0	pSport1	pSport1
H2CBF28	HCQDM28 Lambda ZAP	HUKF071	нсолт28	HCYBC56	HAAAC11	HNHOI84	HRABT72	HWLEG68	99ACIISH	HWAADIS	HUFFD27	HWLMZ30
61	92		06					66	86	83		
53	68		87					66	96	82		
573	434	268	294	427	955	393	364	336	456	507	329	347
1	45	2	<b>1</b>	206	362	148	224	-		-	129	3
5548	5549	5550	5551	5552	5553	5554	5555	5556	5557	5558	5559	5560
gb AAD38411. 1 AF1557	emb CAB\$602 5.1		emb CAB6310					dbj BAA90925 .1	dbj BAA90925 .1	gb AAD41466. 1 AF1028		
(AF155739) axotrophin [Mus musculus] >sp Q9WV66 Q9WV66 AXOTROPHIN. Length = 693	(AL117635) hypothetical protein [Homo sapiens] >emb CAB56025.1  (AL117635) hypothetical protein [Homo sapiens] >pir[T17335[T17335] hypothetical protein DKFZp434G145.1 - human (fragment) >sp CAB56025 CAB56025 Hypothetical 21.0 kd protein (fragment). Length		(AJ242739) mitochondrial tryptophanyl-tRNA synthetase [Homo sapiens] >sp CAB63107 CAB63107 Mitochondrial tryptophanyl-tRNA synthetase precursor (EC 6.1.1.2). Length = 360					(AK000070) unnamed protein product [Homo sapiens] Length = 277	(AK000070) unnamed protein product [Homo sapiens] Length = 277	(AF102851) dolichyl-P-Glc:Man9GlcNAc2-PP-dolichyl glucosyltransferase [Homo sapiens] >sp Q9Y672 Q9Y672 DOLICHYL-P-GLC:MAN9GLCNAC2-PP-DOLICHYL GLUCOSYLTRANSFERASE. Length = 507		the state of the s
875924	875925	875926	875927	875932	875933	875934	875935	875936	875937	875938	875939	875940
1271	1272	1273	1274	1275	1276	1277	1278	1279	1280	1281	1282	1283

HCQCM19 Lambda ZAP	HBWCF70 ZAP Express	pSport1	Lambda ZAP II	HCQDG19 Lambda ZAP	HCQAD16 Lambda ZAP	Lambda ZAP II	HGBBG01 Uni-ZAP XR	pBluescript SK-	Lambda ZAP II	Uni-ZAP XR	Lambda ZAP II	pBluescript SK-	Uni-ZAP XR
НСОСМ19	HBWCF70	HCRON30	HCNAK16	нсорд19	нсодл16	HCQAS16	HGBBG01	HILBF13	нсорп8	HEMGF10	нсорд10	H2CBS17	нетут76
	68						86	98		66			62
	68						86	54		66			51
317	1098	440	168	242	766	220	429	130	314	1289	183	1271	286
126	178	87	85	42	117	89	-	2	99	3	7	981	431
5595	5596	5597	5598	5599	2600	5601	5602	5603	5604	5095	9095	2095	2608
	gb AAA35693. 1		-				gb AAC08966. 1	sp P39193 AL U6_HUMAN		gb AAB61919. 1			gb AAA81016. 1
	cytoplasmic linker protein-170 alpha-2 [Homo sapiens] >pir A43336 A43336 microtubule-vesicle linker CLIP-170 - human Length = 1392						(AF045459) Etk/Bmx cytosolic tyrosine kinase [Homo sapiens] >sp O60564 O60564 ETK/BMX CYTOSOLIC TYROSINE KINASE. Length = 697	iiii ALU SUBFAMILY SP WARNING ENTRY iiii spp3193JAL Length = 593		[Homo sapiens] >sp P78514 P78514 HYPOTHETICAL 48.1 KD PROTEIN (FRAGMENT). Length = 429			novel transcript; similar to transcription factors activation domains; linked at 5" end to AT hook motif of HMGI-C; Method: conceptual translation supplied by author [Homo sapiens] >pir I39058 I39058 hypothetical protein - human (fragment) Length = 70
876022	876023	876024	876025	876026	876027	876028	876029	876030	876034	876039	876044	876045	876048
1318	1319	1320	1321	1322	1323	1324	1325	1326	1327	1328	1329	1330	1331

		<u> </u>	AP	AP	pt	ΑP	Ř	ΑΡ	AP	A.P			A.P			
pSport1	pSport1	pSport1	Lambda ZAP II	Lambda ZAP II	pBluescript SK-	Lambda ZAP II	Uni-ZAP XR	Lambda Z II	Lambda Z II	Lambda Z II	pSport1	pSport1	Lambda ZAP II	pSport1	pSport1	pSport1
HMVBD68	HWLQD17	HCRME16	НСОСП6	HKLAB15	HCYBH57	нсормо8	HSSEA17	HCQDG14 Lambda ZAP	HCQAQ14 Lambda ZAP	HCQBN16 Lambda ZAP	HWLQE13	HWMBS01	HKLAA70	HWLCK07	HISAV29	HWLXE78
87														97	-	
84														76		
276	126	255	348	488	748	111	538	609	451	145	587	181	228	305	390	514
130	1	1	9/	252	92	34	242	409	209	2	267	26	26	E.	172	77
5609	5610	5611	5612	5613	5614	5615	5616	5617	5618	5619	5620	5621	5622	5623	5624	5625
gb AAA73456. 2														gb AAC78645. 1		
beta-galactosidase alpha peptide [Cloning vector pSport1] >sp Q46478 Q46478 BETA-GALACTOSIDASE ALPHA PEPTIDE (FRAGMENT). Length = 113														(AF105424) brush border myosin I [Homo sapiens] >gb AAD31189.1 AF127026_1 (AF127026) brush border myosin I [Homo sapiens] >sp AAC78645 AAC78645 Brush border myosin I. >sp AAD31189 AAD31189 Brush border myosin I. >gb AAA20900.1  myosin [Homo sapiens] {SUB		
876052	876056	876057	876059	876062	876065	876070	816078	876079	876081	876082	980928	840928	680928	876090	876091	876093
1332	1333	1334	1335	1336	1337	1338	1339	1340	1341	1342	1343	1344	1345	1346	1347	1348

HSLHI12 Uni-ZAP XR	Lambda ZAP II	HCQCR12 Lambda ZAP	HPJBW76 Uni-ZAP XR	HCQCD81 Lambda ZAP	pBluescript SK-	HCQCD09 Lambda ZAP	pSport1	HMAKC34 Uni-ZAP XR	HNGBJ13 Uni-ZAP XR	pSport1
HSLHI12	нсосх03	HCQCR12	HPJBW76	нсоср81	HCYBF60	нсосро	HWLVY67	HMAKC34	HNGBJ13	HCFCP28
75			99				59	84		
75			59				45	08		
2	286	1	288	261	356	432	2	383	535	393
178	143	117	211	-	78	91	631	195	272	1
5626	5627	5628	5629	5630	5631	5632	5633	5634	5635	5636
gb AAC74033.			sp P39188 AL U1_HUMAN				dbj BAA84364 .1	emb CAA6380 2.1		
(AE000196) orf, hypothetical protein [Escherichia coli] >pir[B64835]B64835 probable iron-sulfurbinding protein b0947 - Escherichia coli >sp]P75863]P75863 HYPOTHETICAL 40.6 KD PROTEIN. >dbj[BAA35702.1] Hypothetical protein 7.6 [Escherichia coli] {SUB 15-3			!!!! ALU SUBFAMILY J WARNING ENTRY !!!! Length = 591				DEIH-box RNA/DNA helicase [Arabidopsis thaliana] >sp BAA84364 BAA84364 DEIH-box RNA/DNA helicase. Length = 1538	alpha7 nicotinic acetylcholine receptor subunit [Bos taurus] >sp[P54131 ACH7_BOVIN NEURONAL ACETYLCHOLINE RECEPTOR PROTEIN, ALPHA-7 CHAIN PRECURSOR. Length = 499		
876094	876095	876097	840928	876101	876104	876105	876107	876108	876109	876117
1349	1350	1351	1352	1353	1354	1355	1356	1357	1358	1359

	pCMVSport 2.0	HCQDQ31 Lambda ZAP	pCMVSport 3.0	pCMVSport 3.0	pSport1	Uni-ZAP XR	pSport1	HCQBL07 Lambda ZAP	pBluescript SK-	Uni-ZAP XR	pSport1	Lambda ZAP II	pCMVSport 3.0
HCROH40	HKAAK32	нсорозт	ннеем22	HRABR73	HWMBX68	HE8OF49	HWLHY12	нсовгол	H2LAJ32	HSIAD07	HWLNZ56	HLQBA23	НДРQV66
06			99				55						92
06			47				46					•	92
499	507	523	1601	234	523	969	1221	229	169	123	584	408	824
2		338	591	73	329	348	1	86	2	19	267	154	18
5637	5638	5639	5640	5641	5642	5643	5644	5645	5646	5647	5648	5649	2650
emb CAA7488 6.1			emb CAB0770 1.1				emb CAB1678 4.1						dbj BAA90987 .1
DIF-2 protein [Homo sapiens] >emb CAA65304.1  PRG1 [Homo sapiens] >gb AAC33793.1  (AF083421) radiation-inducible immediate early response gene IEX1 [Homo sapiens] >sp P46695 IEX1_HUMAN RADIATION-INDUCIBLE IMMEDIATE-EARLY GENE IEX-1 (IMMEDIATE EARLY PROTEI			similar to Probable rabGAP domains [Caenorhabditis emb CAB0770 elegans] >pir T27026 T27026 hypothetical protein   1.1  Y48E1C.3 - Caenorhabditis elegans >sp O18207 O18207 Y48E1C.3 PROTEIN. Length = 619				putative protein [Arabidopsis thaliana] >sp O23175 O23175 HYPOTHETICAL 52.0 KD PROTEIN. Length = 462						(AK000167) unnamed protein product [Homo sapiens] Length = 463
876118	876121	876123	876126	876127	876137	876139	876140	876141	876142	876146	876151	876152	876153
1360	1361	1362	1363	1364	1365	1366	1367	1368	1369	1370	1371	1372	1373

Uni-ZAP XR	pSport1	Uni-ZAP XR	pCMVSport 3.0	pSport1	ZAP Express	pCMVSport 3.0	pBluescript SK-	pSport1	Uni-ZAP XR	pSport1	pSport1	pBluescript SK-		pCMVSport 3.0	pSport1
HODEJ02	HWMBZ31	HLTCX04	HYABC06	HLYDI04	HBXFF23	HDPBG07	HCYBF02	HTWDI21	HATED01	HWLVU14	HOVCI12	HCYBB01	HCRPM32	HLDNV31	HCRNN03
							74		100	54				87	
							99		97	4				84	
2193	270	226	558	427	520	872	9/	171	345	1229	262	433	296	335	387
1102	-	999	412	308	299	510	2	16	491	564	119	152	102	es .	1
5651	5652	5653	5654	5655	5656	5657	5658	5659	2660	5661	5995	5663	5664	5665	9999
							gb AAB49034. 1		dbj BAA25646 .1	dbj BAA34528 .1				dbj BAA20786 .2	
							alternatively spliced product using exon 13A [Homo sapiens] >splP78525 P78525 MYB PROTO-ONCOGENE PROTEIN (C-MYB). Length = 666		dolichol-phosphate-mannose synthase [Homo sapiens] >emb CAB53749.1  (AL034553) dJ914P20.1 (dolichyl-phosphate mannosyltransferase polypeptide 1, catalytic subunit) [Homo sapiens] >sp O60762 O60762 DOLICHOL-PHOSPHATE-MANNOSE SYNTHASE.	(AB018351) KIAA0808 protein [Homo sapiens] >sp O94900 O94900 KIAA0808 PROTEIN. Length = 526				(AB002326) KIAA0328 protein [Homo sapiens] >sp BAA20786 BAA20786 KIAA0328 protein (fragment). Length = 1906	
876155	876156	876166	876168	876169	876170	876172	876174	876177	876179	876182	876183	876184	876187	876192	876193
1374	1375	1376	1377	1378	1379	1380	1381	1382	1383	1384	1385	1386	1387	1388	1389

HTPIQ89 Uni-ZAP XR	001 pSport1	01 pSport1	C10 pSport1	D53 pCMVSport 3.0	117 pBluescript	12 pSport1	357 pCMVSport 3.0	V09 pSport1	J02 pCMVSport 2.0	pSportl	V14 pSport1	415 pBluescript SK-	J56 Uni-ZAP XR	V93 Uni-ZAP XR
нтріо	HWLQD01	HISAQ01	HCRMC10	HWABD53	HKCSF17	HTDAI12	HYABB57	HWLVN09	HOHAU02	HCRNJ43	HWLGV14	HCYBM15	HTXOU56	HHFCN93
78					100					84	8	8		
78					100			_		83	80	74		
575	296	589	602	711	7	870	433	427	287	1459	826	510	161	995
204	147	80	354	418	298	295	302	152	135	2	46	217	18	834
5667	8995	5669	5670	5671	5672	5673	5674	5675	9299	5677	5678	5679	2680	5681
dbj BAA31678 .1					gb AAB18496.					emb CAA8122 6.1	gb AAD23440. 1 AF1153	gb AAC27698. 1		
(AB014603) KIAA0703 protein [Homo sapiens] >sp 075185 075185 KIAA0703 PROTEIN. Length = 963					No definition line found [Escherichia coli] >gb AAC76545.1  (AE000428) putative regulator [Escherichia coli] >pir S47740 S47740 probable transcription regulator (treF-kdgK intergenic region) - Escherichia coli >sp P37640 YHJB_ECOLI HYPOTHETICAL TRANSCRIPT					desmoglein 2 [Homo sapiens] >pir S38673 S38673 desmoglein 2 - human >sp Q14126 DSG2_HUMAN DESMOGLEIN 2 PRECURSOR (HDGC). Length = 1117	(AF115384) LR8 [Homo sapiens] >sp Q9Y609 Q9Y609 LR8. Length = 270	(AF038388) actin-filament binding protein Frabin [Rattus norvegicus] >sp 088387 088387 ACTIN-FILAMENT BINDING PROTEIN FRABIN. Length = 766		
876198	876200	876201	876206	876207	876208	876209	876213	876215	876220	876224	876226	876228	876229	876232
1390	1391	1392	1393	1394	1395	1396	1397	1398	1399	1400	1401	1402	1403	1404

1405	876236	KIAA0020 [Homo sapiens]   Spi(015397 Y020 HUMAN HYPOTHETICAL   PROTEIN KIAA0020. Length = 508	dbj BAA02808   -1	5682	_	1458	92	92	H2CBC05	pBluescript SK-
1406	876238	S.cerevisiae apiens] ITVE (1 (EC M 1) = 394	dbj BAA07679 .1	5683	2	640	83	87	нтере28	Uni-ZAP XR
1407	876239	(AC004520) similar to NFE2-related transcription factors; similar to 148694 (PID:g2137676) [Homo sapiens] >sp Q9Y4A8 Q9Y4A8 WUGSC:H_RG119C02.1 PROTEIN. >dbj BAA76288.1  (AB010812) NF-E2-related factor 3 [Homo sapiens] {SUB 295-694} Length = 694	gb AAC09039.	5684	1	837	94	95	HUSGL79	pSport1
1408	876259			5895	2	703			HPMFU84	Uni-ZAP XR
1409	876260			9895	260	598			HDLAD09	pCMVSport 2.0
1410	876261			2687	297	530			HCQAW45	HCQAW45 Lambda ZAP
1411	876265	IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	sp Q9Y6Y5 Q9 Y6Y5	5688	3	131	06	93	HCYAC01	pBluescript SK-
1412	876266	(AB024057) vascular Rab-GAP/TBC-containing protein [Homo sapiens] >sp O95759 O95759 VASCULAR RAB-GAP/TBC-CONTAINING PROTEIN. Length = 897	dbj BAA75489 .1	5689	233	81	100	100	HCROF86	pSport1
1413	876269			9699	398	502			H2CBJ83	pBluescript SK-
1414	876270			5691	6	104			H2LAW73	pBluescript SK-
1415	876274			5692	1	222			HWMCL22	pSport1

pSport1	pBluescript SK-	pSport1	pBluescript	HMWFC49 Uni-ZAP XR	Uni-ZAP XR	pSport1	Uni-ZAP XR	pBluescript SK-	HNEDH18 Uni-ZAP XR	pSport1
HCRPZ42	HCYBM32	HCRPJ72	HKCSA58	HMWFC49	HMSIE02	HCRMZ34	HTGAM27	HCYBI20	HNEDH18	нwмғQ61
							82	68		96
			96				82	68		96
338	-	604	178	171	643	366	247	981	327	877
189	441	431	17	-	368	-	2	34	31	2
5693	5694	5695	9695	2692	8698	5699	5700	5701	5702	5703
			dbj BAA14940 .1				emb CAB4671 7.1	emb CAA0066		emb CAA3367
			Thermostable carboxypeptidase (EC 3.4.17). [Escherichia coli] >gb AAC74420.1  (AE000231) putative aminohydrolase (EC 3.5.1.14) [Escherichia coli] >pir E64883 E64883 probable amidohydrolase (EC 3.5) ydaJ - Escherichia coli >sp P77357 YDAJ_ECOLI HYPOT				(AL034396) dJ1158B12.1 (zinc finger, X-linked, duplicated A) [Homo sapiens] >sp CAB46717 CAB46717 DJ1158B12.1 (zinc finger, X-linked, duplicated A). Length = 799	villin [Homo sapiens] >emb CAA31386.1  villin (AA emb CAA0066 1 - 827) [Homo sapiens] >pir A31642 A31642 villin - 4.1  human >sp P09327 VIL1_HUMAN VILLIN. {SUB 2-827} Length = 827		putative precursor (AA 1-304) [Homo sapiens] emb CAA3367 >emb CAA61579.1  uracil-DNA-glycosylase, UNG1   9.1  [Homo sapiens] >pir S05964 A60472 uracil-DNA glycosylase (EC 3.2.2) precursor - human >sp P13051 UNG HUMAN URACIL-DNA GLYCOSYLASE PRECURSOR (EC 3.2.2) (UDG). >e
876276	876277	876278	876280	876281	876282	876284	876300	876304	876306	876308
1416	1417	1418	1419	1420	1421	1422	1423	1424	1425	1426

1427	876309			5704	532	840			HFIUZ10	pSport1
1428	876322	tripeptidyl peptidase II [Homo sapiens] >pir S54376 S54376 tripeptidyl-peptidase II (EC 3.4.14.10) - human >sp P29144 TPP2_HUMAN TRIPEPTIDYL-PEPTIDASE II (EC 3.4.14.10) (TPP II) (TRIPEPTIDYL AMINOPEPTIDASE). Length = 1249	gb AAA36760. 1	5705	٣	521	91	91	HDPJE43	pCMVSport 3.0
1429	876326			90/5	38	268			HWLWR22	pSport1
1430	876327			5707	132	290			HCRNJ16	pSport1
1431	876330			5708	1	192			HPRAZ22	Uni-ZAP XR
1432	876333			5709	133	249			HWLQG81	pSport1
1433	876334			5710	554	688			HOENU48	HOENU48 Uni-ZAP XR
1434	876335	(AK001944) unnamed protein product [Homo sapiens] Length = 526	dbj BAA91992 .1	5711	719	904	87	87	HOUDK26	HOUDK26 Uni-ZAP XR
1435	876340	FAST kinase [Homo sapiens] >pir I37386 I37386 FAST kinase - human >sp Q14296 Q14296 FAST KINASE. Length = 549	emb CAA6044 8.1	5712	84	999	26	86	HODDG78	Uni-ZAP XR
1436	876345	(AK002163) unnamed protein product [Homo sapiens] Length = 642	dbj BAA92116 .1	5713	3	827	86	86	HAMFP80	pCMVSport 3.0
1437	876354			5714	78	359			нwнов10	pCMVSport 3.0
1438	876361			5715	42	182			H2LAB47	pBluescript SK-
1439	876364	(AF092563) chromosome-associated protein-E [Homo sapiens] >spl095347 095347 CHROMOSOME-ASSOCIATED PROTEIN-E. Length = 1197	gb AAC72360. 1	5716	158	370	68	06	HJBAR28	pBluescript SK-
1440	876370	(AB002353) KIAA0355 [Homo sapiens] >sp O15063 Y355_HUMAN HYPOTHETICAL PROTEIN KIAA0355. Length = 1070	dbj BAA20812 .1	5717	162	317	100	100	HCEFA76	Uni-ZAP XR

HCQBI31 Lambda ZAP	HTEGD78 Uni-ZAP XR	HCYBN59 pBluescript SK-	HCYBC31 pBluescript SK-	HCQBM44 Lambda ZAP	HKCSP75 pBluescript	HKCSP84 pBluescript	HPMFF45 Uni-ZAP XR	HE2CT52 Uni-ZAP XR	HTNBJ76 pBluescript SK-	HE9ND38 Uni-ZAP XR	HPIAK40 Uni-ZAP XR	HHPGD10 Uni-ZAP XR	HCQBI47 Lambda ZAP	HE8DW67 Uni-ZAP XR
					-				80		89	100		
									77		57	001		
408	390	142	300	190	258	301	242	459	114	364	538	189	675	321
1	190	2	-	2	145	17	3	307	1	29	729		394	109
5718	5719	5720	5721	5722	5723	5724	5725	5726	5727	5728	5729	5730	5731	5732
						-			sp P39192 AL US_HUMAN		emb CAB6721	dbj BAA76862 .1		
									IIII ALU SUBFAMILY SC WARNING ENTRY IIII sp P39192 AL Length = 585		(AJ271079) hypothetical protein [Oenothera elata subsp. hookeri] >emb CAB67234.1  (AJ271079) hypothetical protein [Oenothera elata subsp. hookeri] >sp CAB67211 CAB67211 Hypothetical 6.7 kd protein. >sp CAB67234 CAB67234 Hypothetical 6.7 kd protein. Length	(AB023235) KIAA1018 protein [Homo sapiens] >sp Q9Y2M0 Q9Y2M0 KIAA1018 PROTEIN. Length = 1017		
876372	876374	876376	876379	876380	876381	876382	876383	876385	876386	876387	876395	876397	876398	876399
1441	1442	1443	1444	1445	1446	1447	1448	1449	1450	1451	1452	1453	1454	1455

pBluescript SK-	ambda ZAP II	pBluescript SK-	pSport1	pBluescript SK-	pBluescript	pSport1	Uni-ZAP XR	pSport1	pBluescript SK-	pBluescript SK-	Uni-ZAP XR	pBluescript SK-	pBluescript	pBluescript SK-	pCMVSport 1	Uni-ZAP XR
HONAH83	HHGCW95 Lambda ZAP	HCYBI75	HCRMK04	H2CBF13	HKCSO44	HWLKU83	HE9RM22 L	HCRPQ93	HPDDL36	Н2СВМ09	HKCAA10 U	H2CB125	HKISB80	H2CBE84	HSEBD08	HPMFM22 U
		84	92			52	86			93					88	
		6/	92			38	86			93					85	
1012	371	484	103	909	107	509	099	116	177	513	375	274	397	156	811	323
758	3	359	2	1	3	405	1	3	-	148	178	86	47		2	123
5733	5734	5735	5736	5737	5738	5739	5740	5741	5742	5743	5744	5745	5746	5747	5748	5749
		gb AAA65999. 1	emb CAB6919 5.1			gb AAB02291. 1	dbj BAA91072 .1			dbj BAA91221 .1					gb AAC60637.	
		neutral protease large subunit [Homo sapiens] Length gb AAA65999. = 166	unnamed protein product [unidentified] Length = 180			reverse transcriptase [Homo sapiens] Length = 361	(AK000307) unnamed protein product [Homo sapiens] Length = 325			(AK000515) unnamed protein product [Homo sapiens] Length = 203					putative cytoskeletal protein=H4(D10S170) [human, gb AAC60637. thyroid, Peptide, 585 aa] [Homo sapiens] >pir I58403 I58403 H4 protein - human >sp Q16204 D170_HUMAN H4(D10S170) . PROTEIN. Length = 585	
876400	876401	876402	876404	876405	876408	876409	876418	876419	876420	876422	876425	876426	876427	876428	876431	876432
1456	1457	1458	1459	1460	1461	1462	1463	1464	1465	1466	1467	1468	1469	1470	1471	1472

pCMVSport 2.0	Uni-ZAP XR	pBluescript SK-	pSport1	pSport1	Uni-ZAP XR	pCMVSport 2.0	pCMVSport 3.0	pCMVSport 3.0	Uni-ZAP XR	pSport1	pCMVSport 3.0		HPWAY46 Uni-ZAP XR	HLTAH77 Uni-ZAP XR	pSport1	pBluescript
HDHEB14	HAIDH43	HJAAL27	HA5AB14	HWLNS47	HE8UJ03	HDTLK03	HMTBC69	HMUBP81	НАРОТ58	HCFLR18	HDPAA38	HCYBM66	HPWAY46	HLTAH77	HWLXX39	HPTWG85
						98	98	100	55							86
						98	85	100	53							97
641	474	424	1017	857	894	1622	693	896	1295	271	483	144	629	507	504	407
186	181	149	754	774	029	3	-	87	ю	95	187	-	327	100	-	6
5750	5751	5752	5753	5754	5755	5756	5757	5758	\$759	2760	5761	5762	5763	5764	5765	5766
						dbj BAA91700 .1	dbj BAA09436 .1	gb AAF36159. 1 AF1510	gb AAF28992. 1 AF1614							emb CAA7662 9.1
						(AK001452) unnamed protein product [Homo sapiens] Length = 718	opeptidase [Homo sapiens] LACENTAL LEUCINE Length = 944	(AF151073) HSPC239 [Homo sapiens] Length = 293 gb AAF36159.	(AF161432) HSPC314 [Homo sapiens] >sp AAF28992 AAF28992 HSPC314 (fragment). Length = 248							rhomboid-related protein [Homo sapiens] >sp 075783 075783 RHOMBOID-RELATED
876435	876436	876440	876441	876444	876447	876448	876451	876452	876458	876459	876464	876465	876469	876470	876471	876472
1473	1474	1475	1476	1477	1478	1479	1480	1481	1482	1483	1484	1485	1486	1487	1488	1489

Uni-ZAP XR	Uni-ZAP XR	pSport1	Uni-ZAP XR	pBluescript SK-	pCMVSport 3.0	pCMVSport 3.0	pCMVSport 2.0
HE6BS09	HERAM35 Uni-ZAP XR	HFIUG54	HE8CX56	H2LAQ54	HWABG32	HMTBE05	HKABL05
29			81	26	16		83
50			81	97	91		83
1054	314	086	573	387	1199	494	1006
2	150	618	-	-	ы	3	455
5767	5768	5769	5770	5771	5772	. 5773	5774
gb AAA68725.			gb AAB84144. 1	gb AAB16860. 1	gb AAA35678.		dbj BAA83340 .1
F08C6.6 gene product [Caenorhabditis elegans] >pir T15973 T15973 hypothetical protein F08C6.6 - Caenorhabditis elegans >sp Q19202 Q19202 COSMID F08C6. Length = 296			(AF029343) protocadherin 68 [Homo sapiens] >pir T09055 T09055 protocadherin 68 - human >sp 014917 014917 PROTOCADHERIN 68. Length = 889	Ptx1 [Mus musculus] >sp P70314 PtX1_MOUSE PITUITARY HOMEOBOX 1 (HOMEOBOX PROTEIN P-OTX) (PITUITARY OTX-RELATED FACTOR) (HINDLIMB EXPRESSED HOMEOBOX PROTEIN BACKFOOT). >gb AAC53059.1  hindlimb expressed homeobox protein backfoot [Mus musculus] {SUB 133-315}	IgG Fc receptor I [Homo sapiens] >gb AAA36049.1  gb AAA35678. Fc gamma receptor I [Homo sapiens] >pir A39878 A39878 Fc gamma (IgG) receptor I-A (high affinity) precursor - human >sp Q92663 Q92663 FC GAMMA RECEPTOR I. Length = 374		(AB030905) Heterochromatin protein I gamma [Homo sapiens] >gb AAB48101.1  HP1Hs-gamma [Homo sapiens] >sp Q13185 CBX3_HUMAN CHROMOBOX PROTEIN HOMOLOG 3 (HETEROCHROMATIN PROTEIN 1 HOMOLOG GAMMA) (HP1 GAMMA) (MODIFIER 2 PROTEIN). >sp BAA83340 BAA83340 Hetero
876473	876474	876475	876476	876480	876481	876483	876484
1490	1491	1492	1493	1494	1495	1496	1497

pSport1	pSport1	pSport1	pCMVSport 3.0	Uni-ZAP XR	pSport1	HAPQU61 Uni-ZAP XR	Uni-ZAP XR	pBluescript SK-
HOCTA74	HWLUU48	HULAJ15	HSYAJ64	HETIF19	HLYEA23	HAPQU61	HE8OT93	H2LAB08
100	55	97	94	79	77		93	100
100	43	26	94	78	89		83	100
292	772	831	1860	1354	148	494	9/1	629
65	602	-	166	836	294	306	465	E.
5775	2776	5777	5778	5779	5780	5781	5782	5783
emb CAA3276 3.1	emb CAA2736 3.1	pir A49800 A4 9800	gb AAC62263. 1	emb CAA6337	dbj BAA91205 .1		gb AAD34062. 1 AF1518	gb AAC31610. 1
propionyl-CoA carboxylase [Homo sapiens] >pir S04613 A27883 propionyl-CoA carboxylase (EC 6.4.1.3) alpha chain precursor - human >sp P05165 PCCA_HUMAN PROPIONYL-COA CARBOXYLASE ALPHA CHAIN PRECURSOR (EC 6.4.1.3) (PCCASE) (PROPANOYL- COA:CARBON DIOXIDE LIGA	ORF 2 (466 aa) [Mus musculus] >sp Q61787 Q61787 ORF 2. Length = 466	galectin 3 (version 2) - human Length = 242	(AC005326) asparagine synthetase [Homo sapiens] >sp Q15666 Q15666 ASPARAGINE SYNTHETASE. Length = 561	UDP-GalNAc:polypeptide N-acetylgalactosaminyltransferase (GalNAc-T3) [Homo sapiens] >sp Q14435 Q14435 Q1D4435 POLYPEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE (EC 2.4.1.41) (PROTEIN-UDPACETYLGALACTOSAMINYLTRANSFERASE) (UDP-GALNAC:POLYPEPTIDE, N-ACETYLGALACTOSAMINYL	(AK000496) unnamed protein product [Homo sapiens] Length = 239		(AF151825) CGI-67 protein [Homo sapiens] >sp Q9Y377 Q9Y377 CGI-67 PROTEIN. Length = 293	(AF081281) lysophospholipase [Homo sapiens] >gb AAD26993.1  (AF077198) lysophospholipase [Homo sapiens] >sp O75608 O75608 LYSOPHOSPHOLIPASE. Length = 230
876487	876490	876491	876494	876495	876496	876498	876499	876503
1498	1499	1500	1501	1502	1503	1504	1505	1506

pSport1	pSport1	Uni-ZAP XR	pSport1	pCMVSport 3.0	HLTAR39 Uni-ZAP XR	pSport1	pSport1	pBluescript	Uni-ZAP XR	pSport1	pSport1	pBluescript	pSport1	pCMVSport 3.0	HRODG74 Uni-ZAP XR	pSportl
HISBB72	HCHBN47	HFADJ29	HWLQP42	HDPAG07	HLTAR39	HWLRF38	HCRNM09	HOBAE30	HATCV09	HCRNE16	HCRPV63	HSKKP02	HOVAN13	HWBEX78	HRODG74	HCROK30
93					<i>L</i> 9							93		100		29
93					09							93		100		39
1106	651	778	579	454	21.6	272	617	669	551	300	368	1058	395	718	200	519
129	1	572	145	116	882	72	<i>L</i> 89	340	240	193	81	399	3	2	3	-
5784	5785	5786	5787	5788	5789	5790	5791	5792	5793	5794	5795	5796	5797	5798	5799	5800
gb AAC50569. 1					sp P39188 AL U1_HUMAN							dbj BAA92110 .1		emb CAB7086 2.1		gb AAD47086. 1 AF1662
LAMP [Homo sapiens] >pir JC4776 JC4776 limbic-system-associated membrane protein precursor-human >sp Q13449 LAMP_HUMAN LIMBIC SYSTEM-ASSOCIATED MEMBRANE PROTEIN PRECURSOR (LSAMP). Length = 338					IIII ALU SUBFAMILY J WARNING ENTRY IIII Length = 591							(AK002149) unnamed protein product [Homo sapiens] Length = 330		(AL137657) hypothetical protein [Homo sapiens] >emb CAB70862.1  (AL137657) hypothetical protein [Homo sapiens] >sp CAB70862 CAB70862 Hypothetical 12.1 kd protein. Length = 106		(AF166261) nuclear protein Sojo [Xenopus laevis] >sp AAD47086 AAD47086 Nuclear protein Sojo. Length = 676
876504	876507	876511	876513	876518	876524	876526	876530	876533	876534	876535	876536	876538	876540	876543	876544	876545
1507	1508	1509	1510	1511	1512	1513	1514	1515	1516	1517	1518	1519	1520	1521	1522	1523

pSportl	pCMVSport 2.0	pSport1	pSport1	Uni-ZAP XR	pCMVSport 2.0	pBluescript SK-	pCMVSport 2.0	pBluescript SK-	HSHAX43 Uni-ZAP XR	pSport1
HDABK73	HOGCO78	HCRNG10	HWLRR08	HTEFP55	HDLAR46	H2CBW66	HOGDS65	H2CBX36	HSHAX43	HCRQI57
92				66			100	88		49
92				66			66	88		35
2508	342	645	165	1102	219	1	836	1043	202	780
352	175	466	1	2	4	375	156	405	2	-
5801	. 5802	5803	5804	5805	9085	5807	5808	5809	5810	5811
dbj BAA24861  -1				gb AAA82605. 1			gb AAD21314. 1	dbj BAA88405		dbj BAA92064 .1
(AB007891) KIAA0431 [Homo sapiens]				CSA protein [Homo sapiens] >pir A57090 A57090 CSA protein - human >sp Q13216 CSA_HUMAN COCKAYNE SYNDROME WD-REPEAT PROTEIN CSA. Length = 396			connexin 26 [Homo sapiens] >pir A43424 A43424 gap junction protein Cx26 - human >sp P29033 CXB2_HUMAN GAP JUNCTION BETA-2 PROTEIN (CONNEXIN 26) (CX26). >sp AD21314 AAD21314 Connexin 26. Length = 226	(AB035443) glycogen-debranching enzyme [Homo sapiens] >dbj BAA88405.1  (AB035443) glycogen-debranching enzyme [Homo sapiens] >sp BAA88405 BAA88405 Glycogen-debranching enzyme (EC 2.4.1.25). Length = 1532		(AK002062) unnamed protein product [Homo sapiens] Length = 469
876546	876548	876549	876551	876553	876557	876558	876559	876560	876572	876575
1524	1525	1526	1527	1528	1529	1530	1531	1532	1533	1534

SK-	HHEGC16 pCMVSport 3.0	H2CBG53 pBluescript SK-	HCYBF23 pBluescript SK-	HODCO80 Uni-ZAP XR	HCYBG67 pBluescript SK-	HCYBI10 pBluescript SK-	H2CBE01 pBluescript SK-	HCYB192 pBluescript SK-	HWMCC28 pSport1
86 HC	нн   66	HZ	72 HC	OH	H	92 HC	90 HZ	92 HC	HW
98	6 66		69			6 06	6 06	06	
1152	1150	297	721	935	370	810	952	166	307
61	320	190	242	723	302	346	2	2	86
5812	5813	5814	5815	5816	5817	5818	5819	5820	5821
gb AAA50598.	gb AAD27712. 1 AF1329		gb AAB60408. 1			gb AAF22800. 1 AF2058	gb AAA82935. 1	gb AAA36478. 1	
soform 2 g of uman	(AF132937) CGI-02 protein [Homo sapiens] >sp Q9Y2Z2 Q9Y2Z2 CGI-02 PROTEIN. Length = 618		AMP deaminase [Homo sapiens] >pir S68146 S68146 AMP deaminase (EC 3.5.4.6), erythrocte, splice form 1a - human >sp AAB60408 AAB60408 AMP deaminase (EC 3.5.4.6). >dbj BAA02240.1  human erythrocyte- specific AMP deaminase [Homo sapiens] {SUB 10-776} >gb AAA5			(AF205889) Axin2 [Mus musculus] >sp AAF22800 AAF22800 Axin2. Length = 840	mitosin [Homo sapiens] >sp Q13246 Q13246 NUCLEAR PHOSPHOPROTEIN MITOSIN. Length = 3113	glycine decarboxylase [Homo sapiens] >pir N0124 N0124 glycine dehydrogenase (decarboxylating) (EC 1.4.4.2) - human >sp P23378 GCSP_HUMAN GLYCINE DEHYDROGENASE [DECARBOXYLATING], MITOCHONDRIAL PRECURSOR (EC 1.4.2.2) (GLYCINE DECARBOXYLASE) (GLYCINE CLEAV	
876576	876579	876580	876581	876583	876588	876589	876591	876592	876595
1535	1536	1537	1538	1539	1540	1541	1542	1543	1544

(AF102542) beta-1,6- acetylglucosamiyltra >gb AAD21525.1  (A) 1,6-N-acetylglucosam [Homo sapiens] >sp O ACETYLGLUCOSAI Length = 438 Length = 438 HEPATITIS C-ASSO AGGREGATE PROT (AK000322) unnamec sapiens] Length = 783 [Homo sapiens] >gb A (AF048686) dTDP-D- [Homo sapiens] >gb A (AF048686) dTDP-gl protein [Homo sapien D-GLUCOSE 4,6-DE >sp AAD50061 AAD	piens]   gb AAD10824.   5822   353   1747   100   HWMAN61   pSport1   ore 4 beta- re 2/4-GnT   3TA-1,6-N-	97 291	5824 294 596 HWMFE48 pSport1	5825 999 1184 HMTBN44 pCMVSport 3.0	5826 2 457 HCROI04 pSport1	gate protein UBULAR 444	Homo dbj BAA91085 5828 2 463 85 88 HETBI79 Uni-ZAP XR	5829 .50 208 HWTBM65 Uni-ZAP XR	5830 536 760 HCQBN77 Lambda ZAP	ratase         emb CAA0684         5831         105         1172         99         99         HKAED74         pCMVSport           18686_1         0.1          2.0           18ce like         5455 DTDP-         2.0           C 4.2.1.46).         0se 4,6-d	
	876596 (AF102542) beta-1,6-N- acetylglucosaminyltransferase [Homo sapiens] >gb AAD21525.1  (AF038650) core 2/core 4 beta- 1,6-N-acetylglucosaminyltransferase; core 2/4-GnT [Homo sapiens] >sp O95395 O95395 BETA-1,6-N- ACETYLGLUCOSAMINYLTRANSFERASE. Length = 438	876597	876600	876601	876602	hepatitis C-associated microtubular aggregate protei p44 [Homo sapiens] >sp Q14496 Q14496 HEPATITIS C-ASSOCIATED MICROTUBULAR AGGREGATE PROTEIN P44. Length = 444	876609 (AK000322) unnamed protein product [Homo sapiens] Length = 783	876610	876612	876621 (AJ006068) dTDP-D-glucose 4,6-dehydratase [Homo sapiens] >gb AAD50061.1 AF048686_1 (AF048686) dTDP-glucose 4,6-dehydratase like protein [Homo sapiens] >sp O95455 O95455 DTDP-D-GLUCOSE 4,6-DEHYDRATASE (EC 4.2.1.46).	876622

pSport1	pSport1	pSport1	Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1
HCRMD40	нгио78	HCRPG35	НЅQFQ92	HUFBF32	HTXCO05	HWMBJ09	HSIDP84	HUSJA29
	62	99		76	83	97		100
	78	64		29	83	97		97
181	762	173	672	1702	681	762	488	388
17	70	3	505	38	_	-	225	95
5833	5834	5835	5836	5837	5838	5839	5840	5841
	pir A40032 A4 0032	gb AAB97620.		dbj BAA22896	gb AAB03694. 1	emb CAA6078 0.1		dbj BAA74900 .1
	transcription enhancer factor TEF1 - human >sp P28347 TEF1_HUMAN TRANSCRIPTIONAL ENHANCER FACTOR TEF-1 (PROTEIN GT-IIC) (TRANSCRIPTION FACTOR 13) (NTEF-1). >gb AAB00791.1  transcription enhancer factor [Homo sapiens] {SUB 16-426} Length = 426	(AC004030) F21856_2 [Homo sapiens] >pir[T00636 T00636 hypothetical protein F21856_2 - human >sp O43360 O43360 F21856_2. Length = 679		hepatoma-derived growth factor [Mus musculus] >pirJJC5662JJC5662 hepatoma-derived growth factor-related protein 2 - mouse >sp[035540]035540 HEPATOMA-DERIVED GROWTH FACTOR, RELATED PROTEIN 2. Length = 669	DNA topoisomerase III [Homo sapiens] >sp Q13472 TP3A_HUMAN DNA TOPOISOMERASE III ALPHA (EC 5.99.1.2). >gb AAB03695.1  DNA topoisomerase III [Homo sapiens] {SUB 26-1001} Length = 1001	Human giant larvae homologue [Homo sapiens] >pir S55474 S55474 Human giant larvae homolog -human >sp Q14521 Q1452 GIANT LARVAE HOMOLOGUE. Length = 1015		(AB020684) KIAA0877 protein [Homo sapiens] >sp O94954 O94954 KIAA0877 PROTEIN (FRAGMENT). Length = 580
876630	876631	876633	876637	876638	876643	876645	876646	876647
1556	1557	1558	1559	1560	1561	1562	1563	1564

876648	(AJ009936) nuclear hormone receptor PRR1-A [Homo sapiens] >sp CAB55489 CAB55489 Nuclear hormone receptor PRR1-A. >emb CAB55491.1  (AJ009936) nuclear hormone receptor PRR1-B [Homo sapiens] {SUB 56-434} Length = 434	emb CAB5548 9.1	5842	8	290	100	100	HCQAG09	HCQAG09 Lambda ZAP
l	cyclin F [Homo sapiens] >sp P41002 CG2F_HUMAN G2/MITOTIC-SPECIFIC CYCLIN F. Length = 786	gb AAB60342.	5843	2	475	79	79	HCROT53	pSport1
	(AF039023) Ran-GTP binding protein; RanBP6 [Homo sapiens] >sp O60518 O60518 RAN-GTP BINDING PROTEIN (FRAGMENT). Length = 1105	gb AAC14260. 1	5844	es .	215	71	74	HOENXS0	HOENXS0 Uni-ZAP XR
	(AC005531) similar to lymphocyte early activation antigen AIM/CD69; similar to Q07108 (PID:g584906) [Homo sapiens] >sp O95043 O95043 WUGSC:H_DJ0701016.2 PROTEIN. Length = 189	gb AAD04729. 1	5845	2	412	100	100	HCEOW20	HCEOW20 Uni-ZAP XR
			5846	2	391			HCRMG16	pSport1
			5847	1	420			НСЕРН79	Uni-ZAP XR
	(AF061795) dynamin-like protein Dymple isoform [Homo sapiens] >sp O60709 O60709 DYNAMIN-LIKE PROTEIN DYMPLE ISOFORM. >sp AAD22412.1  (AF107048) dynamin-like protein variant 4 [Rattus norvegicus] {SUB 448-699} Length = 699	gb AAC35283.	5848	73	654	76	77	HFOYYS6	pSport1
			5849	555	779			HSXDG80	HSXDG80 Uni-ZAP XR
	(AK000294) unnamed protein product [Homo sapiens] Length = 929	dbj BAA91062 .1	5850	2	724	94	94	ннеик77	pCMVSport 3.0

pCMVSport 3.0	Lambda ZAP II	pSport1	Uni-ZAP XR	pBluescript	pCMVSport 2.0	pSport1	pSport1	pSport1	pCMVSport 2.0	pCMVSport 3.0
ННЕБО14	HKIMC75	HWMBI36	HE8TM64	HKLSA57	HOGCV45	HADCX04	HCRPH70	HCRQM22	HKAEB15	HSYAP76
100			93		92	26	96	9/	61	
100 100			91		91	26	96	75	61	
802	377	132	1634	167	1801	126	1660	332	595	353
23	96	466	3	31	2	34	2	3	2	165
5851	5852	5853	5854	5855	5856	5857	5858	5859	5860	5861
gb AAA36657. 1			pir C45439 C4 5439		gb AAF35255. 1 AF1275	gb AAF35255. 1 AF1275	emb CAA4209 9.1	gb AAC51129. 1	gb AAC50462.	
protein serine/threonine kinase [Homo sapiens] >emb CAA54508.1  Cdk-activating kinase [Homo sapiens] >emb CAA55785.1  MO15/CDK-activating kinase (CAK) [Homo sapiens] >emb CAA54793.1  CDK activating kinase [Homo sapiens] >pir A54820 A54820 CDK-activating p			myosin-I, Myr 1b (alternatively spliced) - rat Length = 1107		(AF127577) nuclear factor RIP140 [Homo sapiens] Length = 1158	(AF127577) nuclear factor RIP140 [Homo sapiens] Length = 1158	integrin alpha6 subunit [Homo sapiens] >gb AAB24829.1  integrin subunit alpha 6 [human, Peptide Partial, 102 aa] [Homo sapiens] {SUB 703-804} Length = 1067	hepatocyte nuclear factor-3/fork head homolog 11B [Homo sapiens] Length = 748	PDGF associated protein [Homo sapiens] >gb AAF03506.1 AC004922_3 (AC004922) PDGF associated protein [Homo sapiens] >sp Q13442 HP28_HUMAN 28 KD HEAT- AND ACID-STABLE PHOSPHOPROTEIN (HASPP28) (PDGF ASSOCIATED PROTEIN). >sp AAF03506 AAF03506 PDGF associated	
876677	876680	876683	876685	876687	849928	876690	876693	969928	876697	876701
1574	1575	1576	1577	1578	1579	1580	1581	1582	1583	1584

1585	876716	(AB002357) KIAA0359 [Homo sapiens] >sp O15066[KF3B_HUMAN KINESIN-LIKE PROTEIN KIF3B (MICROTUBULE PLUS END- DIRECTED KINESIN MOTOR 3B) (KIAA0359) (HH0048). Length = 747	dbj BAA20815 .1	5862	8	689	100	100	HCRMV17	pSport1
1586	876719	ome subunit p55 [Homo 523 26s proteasom p55 232 000232 TT P55. Length = 456	dbj BAA19749	5863	554	1447	100	100	ноекс59	HOEKC59 Uni-ZAP XR
1587	876722	(AB001075) galectin-2 related protein [Rattus norvegicus] >sp Q9Z144 Q9Z144 GALECTIN-2 RELATED PROTEIN. Length = 130	dbj BAA74954 .1	5864	-	306	78	92	HKCSL28	pBluescript
1588	876725	nslation (D) [Homo 3164K7.2 r 2, subunit 2	emb CAB4374	5865	120	1154	88	88	HHEFB46	pCMVSport 3.0
1589	876726	(AL133620) hypothetical protein [Homo sapiens] >emb CAB63746.1  (AL133620) hypothetical protein [Homo sapiens] >sp CAB63746 CAB63746 Hypothetical 115.3 kd protein. Length = 1031	emb CAB6374 6.1	5866	233	814	35	53	HWBBS84	pCMVSport 3.0
1590	876728	epiligrin alpha 3 subunit [Homo sapiens] >pir A55347 A55347 adhesive ligand epiligrin, alpha-3 chain form A precursor - human >sp Q16787 LMA3_HUMAN LAMININ ALPHA-3 CHAIN PRECURSOR (EPILIGRIN 170 KD SUBUNIT) (E170). Length = 1713	gb AAA59483.	5867	-	1407	92	92	HSIFZ22	Uni-ZAP XR
1591	876731			8985	1	249			HCRNB80	pSport1
1592	876732			6985	234	584			HTPAY47	Uni-ZAP XR
1593	876743			5870	120	629			H2LBA37	pBluescript SK-
1594	876744			5871	06	809			HWLIP86	pSport1

Jni-ZAP XR	pCMVSport 2.0	Uni-ZAP XR	pBluescript SK-	pSport1	pCMVSport 3.0	Uni-ZAP XR	HHFCP36 Uni-ZAP XR	Uni-ZAP XR
100 HGBAM79 Uni-ZAP XR	HKAFU85	HNFEO67	H2MBA27	HWLMB30	HHEBN60	HOEMQ68 Uni-ZAP XR	ннесрз6	HTXKH86 Uni-ZAP XR
100	29		100		75			62
100	<i>L</i> 9		100		75			94
430	978	174	281	166	643	1261	265	871
35	1	1	E.	2	2	686	110	743
5872	5873	5874	5875	9285	5877	5878	5879	5880
pir S02827 S02 827	gb AAA51773. 1		emb CAA2515		gb AAA16347. 1			gb AAB91536. 1
retinoic acid receptor beta-2 - human >sp P10826 RRB2_HUMAN RETINOIC ACID RECEPTOR BETA-2 (RAR-BETA-2) (RAR-EPSILON). >gb AAD45688.1 AF157483_1 (AF157483) retinoic acid receptor beta 4 [Homo sapiens] {SUB 113-448} Length = 448	amphiregulin [Homo sapiens] >gb AAA51781.1  amphiregulin [Homo sapiens] >pir A34702 A34702 amphiregulin precursor - human >sp P15514 AMPR_HUMAN AMPHIREGULIN PRECURSOR (AR) (COLORECTUM CELL- DERIVED GROWTH FACTOR) (CRDGF). >gb AAA72989.1  synthetic amphireg		pS2 precursor [Homo sapiens] >emb CAA28695.1  emf pS2 [Homo sapiens] >emb CAA36254.1  pS2 protein 5.1  [Homo sapiens] >pir A26667 A26667 pS2 protein precursor - human >sp P04155 PS2 HUMAN PS2 PROTEIN PRECURSOR (HP1.A) (BREAST CANCER ESTROGEN-INDUCIBLE PROTEIN) (PN		splicing factor [Homo sapiens] >sp Q14499 Q14499 gb AAA16347.  SPLICING FACTOR. Length = 530 1			cell cycle protein p38-2G4 homolog [Homo sapiens] >sp O43846 O43846 CELL CYCLE PROTEIN P38-2G4 HOMOLOG. Length = 394
876745	876747	876750	876752	876753	876760	876762	876764	876767
1595	1596	1597	1598	1599	1600	1091	1602	1603

pSport1	pBluescript SK-	Uni-ZAP XR	HUVHP60 Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	pCMVSport 3.0	pSport1	Uni-ZAP XR	HCQAE79 Lambda ZAP
HISCI72	HJACJ75	HTEDS58	ниунр60	HUFCI29	HCRN002	HAUAF56	HHEUM25	HWLQW08	HOEOP07	нсоае <i>79</i>
88	66	66	95	95		88			93	100
88	66	66	95	95		85			93	100
393	629	1505	1208	1140	280	390	933	458	717	808
	2	54	177	46	2	145	628	306	-	2
5881	5882	5883	5884	5885	2886	5887	5888	6885	5890	5891
emb CAA6313 3.1	gb AAB07897.	emb CAA0706	dbj BAA91721 .1	gb AAA51739. 1		gb AAD37371. 1 AF1447			gb AAD42165. 1 AF1050	3932
ORF [Homo sapiens] >sp P51809 SYBL_HUMAN SYNAPTOBREVIN-LIKE PROTEIN 1. Length = 220	DDBb p48 [Homo sapiens] >sp Q92466 Q92466 DDBB P48. Length = 427	A carboxylase [Homo AA07066 Propionyl-CoA 337.1 exon [Homo sapiens] 539	(AK001492) unnamed protein product [Homo sapiens] Length = 706	ing antigen precursor [Homo A27681 nonspecific cross-rsor - human NONSPECIFIC NATIGEN PRECURSOR.		(AF144756) adipocyte lipid-binding protein [Rattus norvegicus] >sp AAD37371 AAD37371 Adipocyte lipid-binding protein. Length = 150			(AF105036) zinc finger transcription factor GKLF [Homo sapiens] >sp AAD42165 AAD42165 Zinc finger transcription factor GKLF. Length = 470	mucin 2 precursor, intestinal - human (fragments) pir A >gb AAA59163.1  mucin [Homo sapiens] {SUB 626-3932 1895} >gb AAA59164.1  MUC2 [Homo sapiens] {SUB 2037-3020} >gb AAA36334.1  intestinal mucin [Homo sapiens] {SUB 1916-2193} >gb AAA59861.1  mucin-like protein [H
876771	876773	876776	876789	876791	876795	876798	876802	876804	876807	876809
1604	1605	1606	1607	1608	1609	1610	1611	1612	1613	1614

1615	876811	(AF034745) LNXp80 [Mus musculus] >pir[T09457 T09457 numb-binding protein LNXp80 - mouse >sp 070263 070263 LIGAND OF NUMB-PROTEIN X (LNXP80). Length = 728	gb AAC40075. 1	5892	-	681	87	93	нсоркэз	HCQDR53 Lambda ZAP
1616	876816			5893	209	472		-	HOEFO36	Uni-ZAP XR
1617	876817	thrombospondin 2 [Homo sapiens] >pir A47379 TSHUP2 thrombospondin 2 precursor - human >sp P35442 TSP2_HUMAN THROMBOSPONDIN 2 PRECURSOR. Length = 1172	gb AAA03703. 	5894	26	1738	98	98	HFIAL22	pSport1
1618	876822			5895	209	783			HWLMN85	pSport1
1619	876823	(AL122042) hypothetical protein [Homo sapiens] >pir[T34520 T34520 hypothetical protein DKFZp564J157.1 - human (fragment) >sp CAB591791CAB59179 Hypothetical 17.9 kd protein (fragment). >emb CAB59179.2  (AL122042) hypothetical protein [Homo sapiens] {SUB 22	emb CAB5917 9.1	5896	2	529	100	100	HCGLC91	pCMVSport 2.0
1620	876829	(AL034423) dJ1185N5.1 (similar to C.elegans Y53C10A.5 protein) [Homo sapiens] Length = 270	emb CAB7686 4.1	5897	722	1336	97	76	HMHBJ66	Uni-ZAP XR
1621	876830	dopa decarboxylase [Homo sapiens] >gb AAA58437.1  aromatic amino acid decarboxylase [Homo sapiens] >pir A33663 DCHUA aromatic-L-amino-acid decarboxylase (EC 4.1.1.28) - human >sp P20711 DCD HUMAN AROMATIC- L-AMINO-ACID DECARBOXYLASE (EC 4.1.1.28) (DOPA DEC	gb AAA20894.	8688	-	288	92	83	HCQDG08	Lambda ZAP II
1622	876831	nidogen [Homo sapiens] Length = 1246	emb CAA5770 9.1	5899	3	296	96	86	HE8BX38	Uni-ZAP XR
1623	876836	PIG-B [Homo sapiens] >pir S71751 S71751 probable dbj BAA07709 GPI mannosyl transferase PIG-B - human   .1    .1    .1    .1	dbj BAA07709 .1	5900	1194	1051	100	100	HMVCR68	pSport1

HFCAI79 Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	pBluescript	pBluescript	pSport1	pSport1	Uni-ZAP XR
HFCAI79	HBIOH43	ноем136	HWHPZ02	HLTAZ90	HHFUM32	HHFAB62	HWLWJ70	HCRPV85	HCE3V58
		100	66	100	82	92		97	
		100	66	100	85	06		96	
573	1217	974	795	356	484	026	229	1398	231
313	696	168	163	126	59	1158	495		31
5901	5902	5903	5904	5905	9069	5907	5908	5909	5910
		gb AAA35736. 1	dbj BAA91622 .1	dbj BAA02656 .1	dbj BAA02420  .1	dbj BAA11486 .1		gb AAC52275.	
		cyclin [Homo sapiens] >gb AAA60040.1  proliferating cell nuclear antigen (PCNA) [Homo sapiens] >pir A27445 WMHUET proliferating cell nuclear antigen - human >sp P12004 PCNA_HUMAN PROLIFERATING CELL NUCLEAR ANTIGEN (PCNA) (CYCLIN). Length = 261	(AK001321) unnamed protein product [Homo sapiens] Length = 209	human homolog of DnaJ protein [Homo sapiens] Length = 397	subunit c precursor [Homo sapiens] 532.1  P1 gene for c subunit of human ATP synthase [Homo sapiens] 704.1  (AL080089) hypothetical o sapiens] >emb CAB45704.1  nypothetical protein [Hom	KIAA0169 protein [Homo sapiens] >sp Q14675 Q14675 KIAA0169 PROTEIN (FRAGMENT). Length = 1745		retinoblastoma-binding protein mRbAp48 [Mus musculus] >pir[149366[149366 G1/S transition control protein-binding protein RbAp48 - mouse >sp Q60972 RB48 MOUSE CHROMATIN ASSEMBLY FACTOR 1 P48 SUBUNIT (CAF-1 P48 SUBUNIT) (RETINOBLASTOMA BINDING PROTEIN P48)	
876837	876842	876856	876858	876865	876866	876870	876873	876876	876878
1624	1625	1626	1627	1628	6791	1630	1631	1632	1633

1634	876882	hnRNP B1 protein [Homo sapiens] >gb AAA60271.1  hnRNP B1 protein [Homo sapiens] >pir A56845 B34504 heterogeneous nuclear ribonucleoprotein B1 - human >sp P22626 ROA2_HUMAN HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEINS A2/B1 (HNRNP A2 AND HNRNP B1). >gb AAD4973	dbj BAA06031 .1	5911	_	636	100	100	HKGBEII	pSport1
1635	876886	(AB032950) KIAA1124 protein [Homo sapiens] >sp BAA86438 BAA86438 KIAA1124 protein (fragment). Length = 1323	dbj BAA86438 .1	5912	437	1591	98	06	HRAEG13	pCMVSport 3.0
1636	876888	putative [Mus musculus] >pir S15785 S15785 heat-stable antigen-related hypothetical protein HSA-C-mouse >sp Q61692 Q61692 PUTATIVE HEAT STABLE ANTIGEN. Length = 141	emb CAA3984 3.1	5913	384	200	36	43	HLIBZ07	pCMVSport 1
1637	876890	DNase protein [Homo sapiens] >gb AAB17022.1  XIB [Homo sapiens] >pirJJC4633JJC4633 DNase I- like endonuclease (EC 3.1) - human >sp P49184 DRNL_HUMAN MUSCLE-SPECIFIC DNASE I-LIKE PRECURSOR (EC 3.1.21) (DNASE X) (XIB). Length = 302	emb CAA6203 7.1	5914	169	1131	94	94	HTPFB46	HTPFB46 Uni-ZAP XR
1638	876892	IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593	sp P39194 AL U7_HUMAN	5915	5	118	72	9/	HDPSS23	pCMVSport 3.0
1639	876901			5916	-	1077			HCEIC29	Uni-ZAP XR
1640	876903			5917	520	292			HE90Y91	Uni-ZAP XR
1641	876904			5918	354	989			HFKFN66	HFKFN66 Uni-ZAP XR
1642	876905	(AF078859) PTD004 [Homo sapiens] >sp Q9Y6G4 Q9Y6G4 PTD004. Length = 396	gb AAD44491.	5919	2	1324	100	100	100 HWMFQ16	pSport1

876909	membrane protein [Homo sapiens] gb AAA59982.  >emb CAA42708.1  MRP-1 (motility related protein) 1  [Homo sapiens] >gb AAA80320.1  CD9 antigen [Homo sapiens] >gb AAC60586.1  CD9 antigen [human, leukocytes, Peptide, 228 aa] [Homo sapiens]  >pir A46123 A40402 CD9 antigen - hu	gb AAA59982.	5920	41	865	88	88	HCRBB01	HCRBB01 Uni-ZAP XR
			5921	281	625			HSAAN15	pBluescript SK-
	CYCLIN-DEPENDENT KINASE (CDK)8 [unidentified] >emb CAA59754.1  CDK8 protein kinase [Homo sapiens] >pir I37227 I37227 cyclin-dependent kinase 8 - human >sp P49336 CDK8_HUMAN CELL_DIVISION PROTEIN KINASE 8 (EC 2.7.1) (PROTEIN KINASE 8 (EC 2.7.1)	emb CAA0358 5.1	5922	134	1006	68	68	HTEKS27	Uni-ZAP XR
ì	mel-13a protein - mouse Length = 132	pir S65785 S65 785	5923	231	530	92	75	HWMBA10	pSport1
	(AF108460) ubinuclein [Homo sapiens] >gb AAF31756.1 AF108461_1 (AF108461) ubinuclein [Homo sapiens] >gb AAA64188.1  VT4 [Homo sapiens] {SUB 348-691} Length = 1134	gb AAF31755. 1 AF1084	5924	-	723	73	98	нсовозя	Lambda ZAP II
	11-beta-hydroxysteroid dehydrogenase type 2 [Homo gb AAC50356. sapiens] >pir S62789 S62789 11beta-hydroxysteroid   1  dehydrogenase (EC 1.1.1.146) type 2 - human Length = 405	gb AAC50356.	5925	2	166	83	83	HWLGQ64	pSport1
			5926	309	572			HCQCV14	HCQCV14 Lambda ZAP
1			5927	8	127			HCR0059	pSport1
			5928	632	026			HCRPN27	pSport1
			5929	357	512			HCRON34	pSport1

HFKFH50 Uni-ZAP XR	pSport1	pSport1	Lambda ZAP II	pCMVSport 3.0	pCMVSport 1	Uni-ZAP XR	Uni-ZAP XR	pBluescript SK-	pSport1	pBluescript SK-
нғкғн50	HCRQG66	HCROW80	HLQER45	HWADQ26	HLJBJ74	HE8TT24	HSSJS63	H2CAA03	HCROI77	H2CBW39
83		100	66			93	86			100
81		90	66			91	96			100
268	256	791	229	218	322	098	969	507	530	444
2	7	327	228	18	2	486	28	250	141	1
5930	5931	5932	5933	5934	5935	5936	5937	5938	5939	5940
gb AAA35956.		gb AAF31171. 1 AF1497	emb CAA4839  4.1			gb AAF29094. 1 AF1614	gb AAA02852.			dbj BAA25502 .1
heparin-binding EGF-like growth factor [Homo sapiens] >gb AAC15470.1  (AC004634) HBGF [Homo sapiens] >pir A38432 A38432 heparin-binding EGF-like growth factor precursor - human >sp Q99075 HBGF_HUMAN HEPARIN-BINDING EGF-LIKE GROWTH FACTOR PRECURSOR (HB-EGF		(AF149770) sentrin/SUMO-specific protease [Homo sapiens] Length = 643	-GI [Homo sapiens] Length =			(AF161479) HSPC130 [Homo sapiens] >sp AAF29094 AAF29094 HSPC130. Length = 473	aminoacylase-1 [Homo sapiens] >dbj BAA03397.1  45kDa protein [Homo sapiens] >dbj BAA03397.1  aminoacylase-1 [Homo sapiens] >pir A47488 A47488 aminoacylase (EC 3.5.1.14) - human >sp Q03154 ACY1_HUMAN AMINOACYLASE-1 (EC 3.5.1.14) (N-ACYL-L-AMINO-ACID AMIDOH			(AB011148) KIAA0576 protein [Homo sapiens] >pir[T00341 T00341 hypothetical protein KIAA0576 - human (fragment) >sp Q9Y4E5 Q9Y4E5 KIAA0576 PROTEIN (FRAGMENT). Length = 1075
876940	876941	876942	876943	876944	876945	876946	876947	876949	876952	876953
1653	1654	1655	1656	1657	1658	1659	1660	1661	1662	1663

M68 pCMVSport 1	F36 pCMVSport 3.0	CE91 pSport1	J36 Uni-ZAP XR	U84 pSport1	IK65 pSport1	Y93 pSport1	3V37 pSport1		M25 pSport1	3V72 pSport1
ННВНМ68	HSYBF36	HWMCE91	HUVFJ36	HLYBU84	HWLMK65	HWLPY93	HWMBV37	HCDME16	HCRQM25	HWMBV72
0 100	63			98		2 76		7 87	3 71	-
100	83			88		75		87	23	
908	1351	147	279	1229	404	881	294	751	590	428
к	909	221	70	8	153	18	-	338	ы	57
5941	5942	5943	5944	5945	5946	5947	5948	5949	5950	5951
gb AAA52373.	dbj BAA75821 .1			sp Q92664 TF3 A_HUMAN		gb AAD29607. 1 AF1142		gb AAD45388. 1 AF1663	dbj BAA31611 .1	
translational initiation factor eIF-2, alpha subunit [Homo sapiens] >sp P05198 IF2A_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 2 ALPHA SUBUNIT (EIF-2- ALPHA). {SUB 2-315} Length = 315	(AB024436) beta-1,4-galactosyltransferase IV [Homo sapiens] >gb AAC39735.1  (AF038662) beta-1,4-galactosyltransferase [Homo sapiens] >gb AAC72493.1  (AF022367) beta-1,4-galactosyltransferase [Homo sapiens] >sp O60513 O60513 BETA-1,4-GALACTOSYLTRANSFERASE.			TRANSCRIPTION FACTOR IIIA (FACTOR A) (TFIIIA). Length = 423		(AF114264) [Homo sapiens] >sp Q9Y2V1 Q9Y2V1 HYPOTHETICAL 53.6 KD PROTEIN. Length = 448		(AF166331) beta crystallin A2 [Homo sapiens] >splQ9Y562 Q9Y562 BETA CRYSTALLIN A2. >emb CAA60148.1  beta A2 crystallin [Homo sapiens] {SUB 158-185} Length = 197	(AB014536) KIAA0636 protein [Homo sapiens] >gb AAD46074.2  (AF077226) copine III [Homo sapiens] >sp O75131 O75131 KIAA0636 PROTEIN. >sp AAD46074 AAD46074 Copine III. Length = 537	
876954	876957	876958	876959	876961	876963	876964	876965	876966	876967	896928
1664	1665	1666	1667	1668	1669	1670	1671	1672	1673	1674

876969			5952	31	417			HCRQK24	pSport1
			5953	293	829			HWLOK80	pSport1
	cDNA EST EMBL:D75703 comes from this gene; cDNA EST yk513g5.3 comes from this gene; cDNA EST yk528b10.3 comes from this gene [Caenorhabditis elegans] >pir[T27134[T27134] hypothetical protein Y53C12B.2 - Caenorhabditis elegans >sp O18216 O18216 Y53C12B.2 PR	emb CAB1649 1.1	5954	2	820	99	84	HNTBD04	pCMVSport 3.0
976976	(AJ001306) PDZ domain protein [Homo sapiens] >sp O15249 O15249 PDZ DOMAIN PROTEIN. Length = 1524	emb CAA0466 6.1	5955	194	469	89	72	HWLUV59	pSport1
876977	(AF125535) pp21 homolog [Homo sapiens] >sp AAF17229 AAF17229 Pp21 homolog. Length = 104	gb AAF17229. 1 AF1255	5956	-	609	100	100	HSUSF13	pBluescript
876978			5957	243	473			H2CBE41	pBluescript SK-
			8565	516	611			HWLFY03	pSport1
			5959	3	170			HE2JX48	Uni-ZAP XR
876983			966	216	461			HNFHD27	HNFHD27 Uni-ZAP XR
876984			5961	109	339			HWLXS11	pSport1
876985	(AF095791) TACC2 protein [Homo sapiens] >sp O95359 O95359 TACC2 PROTEIN (FRAGMENT). Length = 653	gb AAC64968.	2965	1	510	06	06	HCRPG94	pSport1
876987	IMP dehydrogenase (EC 1.1.1.205) I - human -sp P20839 IMD1_HUMAN INOSINE-5"- MONOPHOSPHATE DEHYDROGENASE 1 (EC 1.11.205) (IMP DEHYDROGENASE 1) (IMPDH-I) (IMPD 1). Length = 514	pir A35566 A3 5566	5963	2	166	100	100	HCUG073	HCUGO73 ZAP Express
876989	KIAA0036 [Homo sapiens] >sp Q15051 Y036_HUMAN HYPOTHETICAL PROTEIN KIAA0036. Length = 598	dbj BAA04968 .1	5964	76	1575	83	85	НРМDD49	Uni-ZAP XR
876990			5965	142	282			HCNSF23	pBluescript

1689	876991			9969	146	340			HKDBC15	pCMVSport 1
1690	876992			2962	602	802			HSIGM23	HSIGM23 Uni-ZAP XR
1691	876993			2968	498	698	,		HCQBN43	HCQBN43 Lambda ZAP
1692	876994			6965	306	999			<b>НСQВО03</b>	HCQBO03 Lambda ZAP
1693	876997			5970	335	505			HCQCF85	HCQCF85 Lambda ZAP
1694	876998	(AB020669) KIAA0862 protein [Homo sapiens] dt >sp BAA74885 BAA74885 KIAA0862 protein. 1 Length = 582	dbj BAA74885 .1	5971	291	842	98	88	HUVFS16	HUVFS16 Uni-ZAP XR
1695	877000	reading frame (gag?) [Spleen necrosis virus]  >pir A93904 FOVDA gag polyprotein - avian spleen 3. necrosis virus (fragment) >sp P03342 GAG_AVISN GAG POLYPROTEIN [CONTAINS: CORE PROTEIN P15; INNER COAT PROTEIN P12; CORE SHELL PROTEIN P30] (FRAGMENT). Length =	emb CAA2451 3.1	5972	229	402	50	09	нсовр <i>я</i> 1	HCQBD51 Lambda ZAP
1696	877001	IIII ALU SUBFAMILY SQ WARNING ENTRY IIII sp P39194 AL Length = 593	sp P39194 AL U7_HUMAN	5973	39	143	74	81	HCRMU18	pSport1
1697	877002			5974	1	258			HONAN63	pBluescript SK-
1698	877004			5975	332	490			нсоси65	Lambda ZAP II
1699	877005			9265	107	187			HCRN079	pSport1
1700	900//8			262	364	636			HCRM022	pSport1
1701	877007	(AB014603) KIAA0703 protein [Homo sapiens] dt >sp O75185 O75185 KIAA0703 PROTEIN. Length .1 = 963	dbj BAA31678 .1	8265	2	673	82	82	HFDME46	pSport1
1702	877008			5979	79	543			HCWHN82	HCWHN82 ZAP Express

Uni-ZAP XR	pBluescript	Uni-ZAP XR	Uni-ZAP XR	pSport1	pSport1	pBluescript	Other	Uni-ZAP XR	HOSBX95 Uni-ZAP XR	Uni-ZAP XR	HE9HL05 Uni-ZAP XR	pSport1
<b>ННРЕК</b> 59	HKCTB07	HFPIZ22	HE8FB89	HCRND67	HSPAI01	HOSXA83	HAVTF85	HTEPJ45	HOSBX95	HSIFP30	неэнгоз	HWLMB91
68	100				100	100					88	
88	100				100	100					81	
271	360	325	727	304	303	059	9//	995	522	366	606	316
2	199	38	494	44	-	3	929	81	262	54	259	98
5980	5981	5982	5983	5984		5986	2865	5988	5989	2990	5991	5992
gb AAD42867. 1 AF1551	gb AAF14118. 1 AF1050				gb AAC24984. 1	dbj BAA91218 .1					gb AAA02993. 1	
(AF155101) putative kruppel-related zinc finger protein NY-REN-23 antigen [Homo sapiens] >sp Q9Y5A5 Q9Y5A5 PUTATIVE KRUPPEL-RELATED ZINC FINGER PROTEIN NY-REN-23 ANTIGEN (FRAGMENT). Length = 547	(AF105020) putative protein O-mannosyltransferase [Homo sapiens] >sp AAF14118 AAF14118 Hypothetical 84.2 kd protein. Length = 750				(AF027571) phospholipase C-beta 4 isoform [Rattus norvegicus] >sp O88356 O88356 PHOSPHOLIPASE C-BETA 4 ISOFORM (FRAGMENT). Length = 747	(AK000512) unnamed protein product [Homo sapiens] Length = 335					cytochrome P450 PCN3 [Homo sapiens] >pir A34101 A34101 cytochrome P450 3A5 - human >sp P20815 CP35_HUMAN CYTOCHROME P450 3A5 (EC 1.14.14.1) (CYPIIIA5) (P450-PCN3). >gb AAB00083.1  cytochrome P450 [Homo sapiens] {SUB 1-24} Length = 502	
877009	877010	877011	877012	877013	877014	877015	877018	877019	877020	877022	877023	877024
1703	1704	1705	1706	1707	1708	1709	1710	1711	1712	1713	1714	1715

E	cript -	11	P XR	11	cript -	Ŧ	Έ	P XR	#1	<del>1</del> 1	P XR	E E	11.	P XR	rt]	rt1	11
pSport1	pBluescript SK-	pSport1	Uni-ZAP XR	pSport1	pBluescript SK-	pSportl	pSport1	Uni-ZAP XR	pSport1	pSport1	HTAHC75   Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	pSport1	pSport1	pSport1
HOVEE11	HCYBN69	HWLWN24	HOSOZ37	HCROD37	H2LAF20	HCROD15	HS2SG18	HMCHW12	HWLVS52	HCRPG56	VHC75	HCRPH26	HWLWL67	HOSDU39	HCROS68	HWLRT47	HCRPN44
HO		HWI	HOS	HCF	H2I	HCF	HS	НМС	HM	HCI	HT/	HCI	HM	НО	HCI	HM	HCI
	74				75			66			57						
	74				<i>L</i> 9			66			39						
909	450	287	573	190	629	155	408	909	258	212	1649	326	506	878	265	186	624
403	<i>L</i> 9	132	349	2	2	3	160	168	25	21	369	123	3	288	374	46	175
5993	5994	5995	9665	5997	2998	5999	0009	6001	6002	6003	6004	6005	9009	2009	8009	6009	6010
	B7090				40194.			.65999.			491199						
	emb CAB7090 7.1				gb AAC40194. 1			gb AAA 1			dbj BAA91199 .1						
	(AL137755) hypothetical protein [Homo sapiens] >emb CAB70907.1  (AL137755) hypothetical protein [Homo sapiens] >sp CAB70907 CAB70907 Hypothetical 117.6 kd protein (fragment). Length = 1027				(AF067806) cAMP-specific cyclic nucleotide phosphodiesterase PDE8; MMPDE8 [Mus musculus] >splO88502 CN8A_MOUSE CAMP-SPECIFIC 3",5"-CYCLIC PHOSPHODIESTERASE 8A (EC 3.1.4.17). Length = 823			neutral protease large subunit [Homo sapiens] Length gb AAA65999. = 166			(AK000488) unnamed protein product [Homo sapiens] Length = 427						
		_	_			_	_			_		_					
877025	877026	877027	877029	877030	877031	877032	877034	877037	877043	877044	877046	877047	877049	877050	877051	877052	877056
1716	1717	1718	1719	1720	1721	1722	1723	1724	1725	1726	1727	1728	1729	1730	1731	1732	1733

345       337     38     54       425     1       294     1       95     1       484     1       367     1       135     100       105     100       193     1       440     1
gb AAA16358. 6013 2 337 38 54  1
6015 141 425 6015 1 6015 1 87 6015 1 87 6016 136 294 617 9 95 6018 251 484 619 6020 186 371 6020 186 371 6022 7 135 6022 7 135 6024 2 250 6025 2 193 6025 2 193 6025 306 832 1 6017 1 102
6015 1 87
6016 136 294   1
6017 9 95   Feb
6018   251   484
6019     191     367       6020     186     371       6021     2     649       6022     7     135       2      165     100       6024     2     250       6025     2     193       6026     506     832       6027     3     440
6020   186   371
gb AAA73456. 6023
gb AAA73456. 6023
gb AAA73456. 6023 1 165 100 100 1 2  6024 2 250   6025 2 193   6026 506 832   100 100 100 100 100 100 100 100 100 100
2 250 2 193 506 832 3 440
2 193 506 832 3 440
506     832       3     440
3 440
6028 105 770 HTLGE26

pSport1	Uni-ZAP XR	Uni-ZAP XR	pSport1	Lambda ZAP II	pSport1	Uni-ZAP XR	pCMVSport 3.0	pCMVSport 3.0	Uni-ZAP XR	pSport1	pSport1	pSport1	pSport1	HPWBM91 Uni-ZAP XR	pSport1	pCMVSport 3.0	pSport1
HCFDE85	нғеан85	HE8QT45	HWLQL84	нсось82	HCRMW80	HSIGL73	ннеут40	нроно51	HODGR31	HWLWB92	HWLRD79	HWLOW72	HUSGT72	HPWBM91	HWLVB03	HAJAM74	HHMME78
		_	95		49			73		_				96			
			33		39			73						96			
305	221	166	442	210	488	470	592	1168	580	303	235	286	765	453	373	451	486
159	3	2	2	-	3	279	338	527	386	139	74	99	112	136	173	218	-
6029	6030	6031	6032	6033	6034	6035	9809	6037	6038	6039	6040	6041	6042	6043	6044	6045	6046
			gb AAA68258. 1		gb AAF01517. 1 AC0099			dbj BAA91918 .1						gb AAD52585. 1 AF1723			
			similar to beta-transducin [Caenorhabditis elegans] >pir T16607 T16607 hypothetical protein K10B2.1 - Caenorhabditis elegans >sp Q09990 YSS1_CAEEL HYPOTHETICAL 80.3 KD TRP-ASP REPEATS CONTAINING PROTEIN K10B2.1 IN CHROMOSOME II. Length = 701		(AC009991) protein [Arabidopsis thaliana] >sp AAF01517 AAF01517 F9F8.14 protein. Length = 701			(AK001798) unnamed protein product [Homo sapiens] Length = 298						(AF172328) [Homo sapiens] >sp AAD52585 AAD52585 Hypothetical 10.2 kd protein. Length = 95			
877092	877093	877094	877095	877096	877097	877098	877099	877101	877104	877105	877106	877110	877111	877112	877114	877119	877120
1752	1753	1754	1755	1756	1757	1758	1759	1760	1761	1762	1763	1764	1765	1766	1767	1768	1769

pBluescript SK-	pSport1	pSport1	pSport1	Uni-ZAP XR	HCQAK62 Lambda ZAP	HCQDP71 Lambda ZAP	Uni-ZAP XR	Lambda ZAP II	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pBluescript SK-	pSport1	pBluescript	nSport1
HCYBJ73	HCRNE77	HWMBC94	HWLMS73	HFAMB70	НСQАК62	нсорр71	HE9PB28	HCQCR68	HEPNB10	HWLNY36	HWLRC68	HWLQM88	HWLMG40	HWLQ015	H2CAC59	HWLXJ87	HSDSJ26	HCFRR55
									86	78	100				98			
									96	73	100				84			
2	465	332	982	662	291	277	1513	298	167	238	548	309	472	153	220	433	447	550
145	244	201	601	486	139	11	1238	155	3	71	210	178	296	1	2	257	250	413
6047	6048	6046	0509	6051	6052	6053	6054	6055	9509	6057	6058	6029	0909	6061	6062	6063	6064	5909
									emb CAA5805 8.1	sp P39194 AL U7_HUMAN	gb AAF36524. 1 AF1320				\$P Q9Y6Y5 Q9 Y6Y5			
									sodium-D-glucose cotransporter [Homo sapiens] >sp Q92681 Q92681 SODIUM-D-GLUCOSE COTRANSPORTER. Length = 617	IIII ALU SUBFAMILY SQ WARNING ENTRY   IIII   sp P39194 AL    Length = 593	(AF132021) myosin X [Homo sapiens] >gb AAF17363.1 AF184153_1 (AF184153) myosin X [Homo sapiens] {SUB 347-495} Length = 1540				IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414			
877121	877122	877123	877126	877129	877130	877131	877132	877133	877134	877135	877137	877138	877139	877140	877142	877143	877145	877146
1770	1771	1772	1773	1774	1775	1776	1777	1778	1779	1780	1781	1782	1783	1784	1785	1786	1787	1788

	T	~	T	l	~	1	٠		۵		а	<u>~</u>		
pSport1	pSport1	Uni-ZAP XR	pCMVSport 2.0	pBluescript SK-	Uni-ZAP XR	pBluescript	pCMVSport 2.0	pSport1	HCQCT53 Lambda ZAP	pSport1	HCQDP52 Lambda ZAP	HFAAH06 Uni-ZAP XR	pSport1	pBluescript SK-
HCRNP62	HCRMR04	нсвне60	HKAOG63	H2CBR38	HRDEW54	HBMDC60	HOGDM40	HWLNG61	<b>НСQCT53</b>	HCRNV59	нсорр52	HFAAH06	HWLMX02	HCYBH52
89										92			09	
41										82			09	
409	318	710	263	531	154	219	712	307	439	219	362	694	606	468
2	16	453	18	382	2	64	407	197	549	1	216	581	-	286
9909	2909	8909	6909	0209	6071	6072	6073	6074	6075	9209	2209	8209	6209	0809
emb CAA9033 8.1										dbj BAA92068 .1			gb AAD42872. 1 AF1551	
cDNA EST yk321h8.5 comes from this gene; cDNA EST EMBL:D68896 comes from this gene; cDNA EST yk395f9.5 comes from this gene; cDNA EST yk360f12.5 comes from this gene [Caenorhabditis elegans] >emb CAA21522.1  (AL032624) cDNA EST yk321h8.5 comes from this g										(AK002071) unnamed protein product [Homo sapiens] Length = 528			(AF155106) NY-REN-36 antigen [Homo sapiens] >sp Q9Y5A1 Q9Y5A1 NY-REN-36 ANTIGEN (FRAGMENT). Length = 227	
877147	877148	877149	877153	877154	877155	877157	877163	877165	877166	877167	877168	877169	877170	877171
1789	1790	1791	1792	1793	1794	1795	1796	1797	1798	1799	1800	1801	1802	1803

pSport1	pCMVSport 3.0	HCQAB45 Lambda ZAP	pBluescript SK-	HCQDF43 Lambda ZAP	HSHBU44 Uni-ZAP XR	pBluescript	HOSDV69 Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1	pCMVSport 3.0	pSport1
HCRNX51	ННЕРР92	HCQAB45	HCYBG53	HCQDF43	HSHBU44	HLHSE50		HCRMH42	HSKZE25	HCRMP38	HDPXD55	HHMMB40
78							100			91		
78							76			16		
231	148	158	281	383	856	162	423	396	1012	507	347	174
	279	ω	48	183	674	13	-	202	758	238	213	28
6081	6082	6083	6084	6085	9809	2809	8809	6809	0609	6091	6092	6093
dbj BAA75500 .1							dbjBAA28346 .1			emb CAB6306		
(AB018122) FGF-19 [Homo sapiens] >gb AAD45973.1 AF110400_1 (AF110400) fibroblast growth factor 19 [Homo sapiens] >sp O95750 O95750 FGF-19. >sp AAD45973 AAD45973 Fibroblast growth factor 19. Length = 216							(AB008164) ST1C2 [Homo sapiens] >gb AAC51285.1  sulfotransferase [Homo sapiens] >gb AAC00409.1  (AF026303) sulfotransferase [Homo sapiens] >sp O00338 O00338 SULFOTRANSFERASE. Length = 296			(AL023654) dJ549K18.1 (novel protein similar to GS2) [Homo sapiens] >sp CAB63061 CAB63061 DJ549K18.1 (novel protein similar to GS2) (fragment). Length = 326		
877173	877174	877175	877176	877181	877184	877185	877187	877189	877191	877194	877195	877200
1804	1805	1806	1807	1808	1809	1810	1811	1812	1813	1814	1815	1816

pCMVSport 3.0	pBluescript	pSport1	pSport1	pSport1	pBluescript	pCMVSport 3.0	pBluescript SK-	Lambda ZAP II	pSport1	pSport1	pSport1	Lambda ZAP II	pCMVSport 3.0	pSport1	pSport1	pSport1
HEQAN41	HSDZB30	HWLWH56	HWLOT46	HOVCR67	HLHSV54	HSYBZ84	H2LAC34	HCQAE29	HCRMV19	HWLMF31	HFIIZ28	нсорк28	ННЕQ129	HTWFA44	HOCMF20	HWMB050
82	74				75		09								100	
77	72				59		43								100	
175	289	256	314	143	496	969	502	537	218	429	195	151	288	503	366	391
2	23	161	120	3	2	514	35	343	3	19	1	2	88	363	1	146
6094	\$609	9609	2609	8609	6609	6100	6101	6102	6103	6104	6105	6106	6107	6108	6109	6110
gb AAF24046. 1 AF0909	gb AAC31117. 1				dbj BAA83010 .1		emb CAA2089 1.1 .								emb CAA7114 3.1	
(AF090931) PRO0483 [Homo sapiens] >sp AAF24046 AAF24046 PRO0483. Length = 60	Ki-1/57 intracellular antigen [Homo sapiens] >spl075804 075804 KI-1/57 INTRACELLULAR ANTIGEN (FRAGMENT). Length = 299				(AB028981) KIAA1058 protein [Homo sapiens] >sp BAA83010 BAA83010 KIAA1058 protein (fragment). Length = 1534		(AL031581) /prediction=(method:"genscan", version:"1.0", score:"198.31"); /prediction=(method:"genefinder", version:"084"); /motif=(desc:"Endoplasmic reticulum targeting sequence", dbase:"PROSITE", acc:"PS00014", method:"ppsearch"); />>								high mobility group protein 2a [Homo sapiens] >sp 015347 015347 HIGH MOBILITY GROUP PROTEIN 2A. Length = 200	
877202	877205	877206	877207	877208	877211	877212	877213	877214	877215	877218	877220	877222	877229	877230	877231	877232
1817	1818	1819	1820	1821	1822	1823	1824	1825	1826	1827	1828	1829	1830	1831	1832	1833

01/23			61111	196	297			НСОВD64	HCQBD64 Lambda ZAP
77234			6112	417	229			HATAP30	HATAP30 Uni-ZAP XR
77235	7 TB1	b AAA03587.	6113	-	759	93	93	H2LBB51	pBluescript SK-
77237			6114	327	830			H6EDT19	Uni-ZAP XR
377240	(AF180919) RNA lariat debranching enzyme [Homo gl sapiens] >sp[AAD53327]AAD53327 RNA lariat debranching enzyme. Length = 544	b AAD53327.	6115	3	542	86	86	HWLOW87	pSport1
377242			6116	176	364			HWLMB22	pSport1
377247	IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  sq (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	p Q9Y6Y5 Q9	6117	£	218	88	88	H2CBA14	pBluescript SK-
877250	(AF234783) tescalcin [Mus musculus] Length = 214 gl	b AAF40439.	6118	-	171	96	86	HCRNM80	pSport1
877251			6119	76	357			НСОСС04	HCQCC04 Lambda ZAP
877254	heat-stable enterotoxin receptor [Homo sapiens]  >pir A40940 OYHUHX heat-stable enterotoxin receptor precursor - human >sp P25092 HSER_HUMAN HEAT-STABLE ENTEROTOXIN RECEPTOR PRECURSOR (GC-C) (INTESTINAL GUANYLATE CYCLASE) (EC-4.6.1.2) (STA RECEPTOR). Lengt	b AAA36655.	6120	14	400	92	96	HCQCI17	HCQCI17 Lambda ZAP
877255			6121	109	324			HFIYJ63	pSport1
877256			6122	379	480			HWLOW51	pSport1
877257			6123	135	341			HHFBA07	Uni-ZAP XR
877258			6124	77	316			HWLD051	pSport1
877261			6125	3	278			HLSAE05	pSport1
877263			6126	3	317			HCRP105	pSport1
	877234 877234 877237 877240 877242 877254 877254 877255 877256 877256 877256 877256 877256	TB1 [Homo sapiens] >sp Q04197 Q04197 TB1 PROTEIN (FRAGMENT). Length = 434  (AF180919) RNA lariat debranching enzyme [Homo sapiens] >sp AAD53327 AAD53327 RNA lariat debranching enzyme. Length = 544  IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414  (AF234783) tescalcin [Mus musculus] Length = 214  heat-stable enterotoxin receptor [Homo sapiens] >pir A40940 OYHUHX heat-stable enterotoxin receptor precursor - human >sp P25092 HSER_HUMAN HEAT-STABLE ENTEROTOXIN RECEPTOR PRECURSOR (GC-C) (INTESTINAL GUANYLATE CYCLASE) (EC-4.6.1.2) (STA RECEPTOR). Lengt	TB1 [Homo sapiens] >sp Q04197 Q04197 TB1 PROTEIN (FRAGMENT). Length = 434  (AF180919) RNA lariat debranching enzyme [Homo sapiens] >sp AAD53327 AAD53327 RNA lariat debranching enzyme. Length = 544  IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414  (AF234783) tescalcin [Mus musculus] Length = 214  heat-stable enterotoxin receptor [Homo sapiens] >pir A40940 OYHUHX heat-stable enterotoxin receptor precursor - human >sp P25092 HSER_HUMAN HEAT-STABLE ENTEROTOXIN RECEPTOR PRECURSOR (GC-C) (INTESTINAL GUANYLATE CYCLASE) (EC-4.6.1.2) (STA RECEPTOR). Lengt	TB1 [Homo sapiens] >sp Q04197 Q04197 TB1 gb AAA03387.  PROTEIN (FRAGMENT). Length = 434 1   (AF180919) RNA lariat debranching enzyme [Homo gb AAD53327. sapiens] >sp AAD53327 AAD53327 RNA lariat debranching enzyme. Length = 544  [DN4-GGTR14 PROTEIN. >db BBAA77334.1  sp Q9Y6Y5 Q9 (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 159-414}    [AF234783] tescalcin [Mus musculus] Length = 214 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 214 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 214 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 214 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 216 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 216 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 216 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 217 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 218 gb AAF40439.    [AF234783] tescalcin [Mus musculus] Length = 218 gb AAF40439.    [AF24414]	TB1 [Homo sapiens] >sp Q04197 Q04197 TB1	TB1 [Homo sapiens] >= p[Q04197[Q04197 TB1]   Bb AAA03587.   6112   417     PROTEIN (FRAGMENT). Length = 434   1   6114   327     (AF180919) RNA lariat debranching enzyme [Homo gb AAD53327.   6115   3     sapiens] >= pp[AAD53327]AAD53327 RNA lariat   2   6116   176     IDN4-GGTR14 PROTEIN. >= db]BAA77334.1   sp[Q9Y6V5]Q9   6117   3     AB019493) IDN4-GGTR9 [Homo sapiens] {SUB   S9-414}   sp[Q9Y6V5]Q9   6117   3     Length = 414   (AF234783) tescalcin [Mus musculus] Length = 214   gb AAF40439.   6118   1     heat-stable enterotoxin receptor [Homo sapiens] gb AAA36655.   6120   41     heat-stable enterotoxin receptor [Homo sapiens] gb AAA36655.   6120   41     PROTEIN AL GIANNYLATE CYCLASE) (GC-C) (INTESTINAL GIANNYLATE CYCLASE) (GC-C) (INTESTINAL GIANNYLATE CYCLASE) (EC-C) (INTESTINAL GI	TB1 [Homo sapiens] >sp[Q04197[Q04197 TB1]   Bb AAA03587.   6112   417   677	TB1 [Homo sapiens] >sp[Q04197[Q04197 TB1]   gb AAA03387, 6113   1   759   93   PROTEIN (FRAGMENT). Length = 434   1   6114   327   830   6114   327   830   6114   327   830   6114   327   830   6114   327   830   6114   327   830   6114   327   830   6114   327   830   6114   327   830   6114   327   830   6114   327   830   6480194919   Evaptical Evaptical Evaptical Evaptical State   544   6116   176   364   648019493)   IDN4-GGTR14 PROTEIN >db BAA77334.1    sp Q3Y6X5[Q9   6117   3   218   85   7414   >emb(CAA2208.1   Homo sapiens] {SUB 159414}   6119   76   357   6119   76   357   6114   647234783) rescalcin [Mus musculus] Length = 214   gb AAA36655.   6120   41   400   92   6117   400   92   6117   619   6118   6119   76   357   6118   6118   6119   76   357   6118   611	TBI [Homo sapiens] >sp QO4197(Q04197 TB1   gb AAA03587   6113   1   759   93   93    RROTEIN (FRAGMENT). Length = 434   1   6114   327   830   830   645   6

pBluescript SK-	pBluescript	pSport1	pSport1	Lambda ZAP II	Lambda ZAP II	HLHE146 Uni-ZAP XR	pSport1	Uni-ZAP XR
HCYBD05	HKLSD44	HFIXP45	HAQNS64	нсордо	нсосрв1	HLHE146	HCROB02	HFKIN68
62	79	94		63	61	86		901
62	78	94		42	57	97		100
484	534	1470	535	250	308	541	346	637
359	7	856	347	2	150	2	221	2
6127	6128	6129	6130	6131	6132	6133	6134	6135
sp P39188 AL U1_HUMAN	gb AAA36433. 1	dbj BAA19968 .1		emb CAB0299 4.2	gb AAC25457.	gb AAA35652. 1		emb CAA4141 8.1
!!!! ALU SUBFAMILY J WARNING ENTRY !!!!  Length = 591	peptide YY [Homo sapiens] >dbj BAA02997.1  peptide YY precursor [Homo sapiens] >pir S33795 S33795 peptide YY (clone S) - human >sp P10082 PYY HUMAN PEPTIDE YY PRECURSOR (PYY). >dbj BAA02998.1  peptide YY precursor variant [Homo sapiens] {SUB 1-90} >pir A3	BMAL1a [Homo sapiens] >sp O00327 BMAL_HUMAN BMAL1 PROTEIN (BRAIN AND MUSCLE ARNT-LIKE 1) (MEMBER OF PAS PROTEIN 3) (MOP3) (BHLH-PAS PROTEIN JAP3). Length = 583		cDNA EST yk552d5.3 comes from this gene [Caenorhabditis elegans] >pir[T21378[T21378] hypothetical protein F25H9.7 - Caenorhabditis elegans >sp[CAB02994 CAB02994 F25H9.7 protein. Length = 137	(AF014898) NADH dehydrogenase subunit 2 [Homo gb AAC25457. sapiens] >sp AAC25457 AAC25457 NADH dehydrogenase subunit 2 (fragment). Length = 347	mast cell carboxypeptidase A precursor [Homo sapiens] >gb AAA59568.1  carboxypeptidase A [Homo sapiens] >pir A43929 A43929 carboxypeptidase A (EC 3.4.17.1) CPA3 precursor -human >sp P15088 CBPC_HUMAN MAST CELL CARBOXYPEPTIDASE A PRECURSOR (EC 3.4.17.1) (		laminin A chain [Homo sapiens] Length = 2628
877264	877272	877274	877275	877280	877281	877282	877283	877284
1850	1851	1852	1853	1854	1855	1856	1857	1858

pCMVSport 3.0	pBluescript SK-	pBluescript SK-	HCE2C40 Uni-ZAP XR	HMCDH54 Uni-ZAP XR	Uni-ZAP XR	pBluescript SK-	HCQAD77 Lambda ZAP	pBluescript	pBluescript	pSport1	Lambda ZAP II	Uni-ZAP XR
HWHGC93	H2CBC75	H2LAW79	HCE2C40	HMCDH54	HTPFG64	Н2СВQ45	HCQAD77	HKLSB60	нгнтс92	HWLXP93	HUKBC55	НЕ9ҒН60
	88			88	100				86			
	98			88	100				76			
655	558	263	264	1784	1133	307	130	363	478	225	86	127
431	1	3	7	3	က	47	2	136	143	-	3	2
6136	6137	6138	6139	6140	6141	6142	6143	6144	6145	6146	6147	6148
	dbj BAA13195 .1			dbj BAA86483 .1	gb AAA36338. 1				gb AAB47250. 1			
	KIAA0204 protein [Homo sapiens] >sp Q92603 Q92603 KIAA0204 PROTEIN. Length = 1152			(AB032995) KIAA1169 protein [Homo sapiens] >sp BAA86483 BAA86483 KIAA1169 protein (fragment). Length = 775					neuronal PAS2 [Homo sapiens] >sp Q99743 NPA2_HUMAN NEURONAL PAS DOMAIN PROTEIN 2 (NEURONAL PAS2) (MEMBER OF PAS PROTEIN 4) (MOP4). Length = 824			
877285	877287	877288	877289	877290	877295	877298	877299	877301	877310	877319	877320	877321
1859	1860	1861	1862	1863	1864	1865	1866	1867	1868	1869	1870	1871

pCMVSport 3.0	HCEOF08 Uni-ZAP XR	Uni-ZAP XR	pSport1	pSport1	pSport1	pBluescript SK-	pBluescript SK-	pSport1	pBluescript SK-	pBluescript SK-
ннеғс89		HLHBZ17	HWLRP86	HISEQ81	HWLWA07	H2CBS31	H2CBN88	HWLOK01	H2CBR23	HCYBK82
	100			98			9		93	86
	100			84			29		93	86
222	105	442	198	241	594	493	444	246	1162	379
-	şd	305	1	396	346	206	178	139	2	137
6149	6150	6151	6152	6153	6154	6155	6156	6157	6158	6159
	gb AAB87524. 1			gb AAB66528. 1			emb CAA6322 4.1		gb AAF18307. 1 AF1328	gb AAF37005. 1
	(AF034374) molybdenum cofactor biosynthesis protein C [Homo sapiens] >sp 014941 014941 MOLYBDENUM COFACTOR BIOSYNTHESIS PROTEIN C. >emb CAA11898.1  (AJ224328) MOCSIB protein [Homo sapiens] {SUB 27-249} Length = 249			(AF009668) polyprotein [multiple sclerosis associated retrovirus] >sp O36581 O36581 POLYPROTEIN (FRAGMENT). >gb AAB66527.1  (AF009666) protease [multiple sclerosis associated retrovirus] {SUB 1-114} Length = 768			STM-7 [Homo sapiens] >sp Q92749 Q92749 TYPE I emb CAA6322 PHOSPHATIDYLINOSITOL-4-PHOSPHATE 5- 4.1  KINASE BETA (EC 2.7.1.68) (STM-7 PROTEIN). >gb AAC50916.1  type I phosphatidylinositol-4-phosphate 5-kinase beta [Homo sapiens] {SUB 112-502} >gb AAC50914.1  type I phosphati		(AF132818) colon Kruppel-like factor [Homo sapiens] >sp AAF18307 AAF18307 Colon Kruppel-like factor. >dbj BAA03393.1  BTEB2 [Homo sapiens] {SUB 239-457} Length = 457	(AF131882) basic-transcription-element-binding- protein 2 [Oryctolagus cuniculus] Length = 219
877324	877326	877327	877329	877331	877332	877333	877334	877336	877338	877339
1872	1873	1874	1875	1876	1877	1878	1879	1880	1881	1882

pSport1	pCMVSport 2.0	Uni-ZAP XR	pCMVSport 2.0	pBluescript SK-	Lambda ZAP II	Lambda ZAP II	Uni-ZAP XR
HCRMK82	HDTB006	HEGAM94 Uni-ZAP XR	HDTAH72	HARAG42	HCQDL20 Lambda ZAP II	HLQGF34	HCDCF78
100				06	100	100	100
100				06	100	100	100
427	765	1693	1515	316	305	514	359
2	559	1406	1357	2	66	278	3
6160	6161	6162	6163	6164	6165	6166	6167
gb AAA51598.				gb AAC05124. 1	gb AAA02993.	gb AAA02993.	dbj BAA04959
adenosine A2b receptor [Homo sapiens] >emb CAA48505.1  A2b adenosine receptor [Homo sapiens] >pir JC1229 JC1229 adenosine receptor A2b - human >sp P29275 AA2B_HUMAN ADENOSINE A2B RECEPTOR. Length = 332				(AF048700) gastrointestinal peptide [Homo sapiens] gb AAC05124. >sp O60575 O60575 GASTROINTESTINAL 1  PEPTIDE. Length = 86	cytochrome P450 PCN3 [Homo sapiens] gb AAA02993.   SpirlA34101 A34101 cytochrome P450 3A5 - human   SpirlA34101 A34101 cytochrome P450 3A5 - human   SpirlA34101 cytochrome P450 PCN3).   SpirlA3400083.1  cytochrome P450 [Homo sapiens]   {SUB 1-24} Length = 502	cytochrome P450 PCN3 [Homo sapiens] >pir A34101 A34101 cytochrome P450 3A5 - human >sp P20815 CP35_HUMAN CYTOCHROME P450 3A5 (EC 1.14.14.1) (CYPIIIA5) (P450-PCN3). >gb AAB00083.1  cytochrome P450 [Homo sapiens] {SUB 1-24} Length = 502	6-pyruvoyl-tetrahydropterin synthase [Homo sapiens] >dbj BAA04224.1  6-pyruvoyl-tetrahydropterin synthase [Homo sapiens] >gb AAA51541.1  6-pyruvoyltetrahydropterin synthase [Homo sapiens] >gb AAB64229.1  putative [Homo sapiens] >gb AAB64229.1  futative [Homo sapiens] >gb AAC16970.1  6-pyruvoyl-
877340	877344	877346	877347	877351	877355	877356	877358
1883	1884	1885	1886	1887	1888	1889	1890

HMIBE59 Uni-ZAP XR	pSport1	Uni-ZAP XR	Uni-ZAP XR	pCMVSport	pBluescript
HMIBE59	HMKAK86	H6EDF71	HOELC15	HAJBN08	<b>Н</b> FVHT62
100		94	06	95	91
100		94	06	95	91
577	293	1059	1081	539	717
2	141	_	∞	120	31
6168	6919	6170	6171	6172	6173
gb AAA36547.		emb CAA4919 6.1	gb AAA52541.	emb CAA0524	gb AAA60222.
ras-like protein [Homo sapiens] >pir D34788 TVHUC4 transforming protein ras (teratocarcinoma clone TC10) - human >sp P17081 RTC0_HUMAN GTP-BINDING PROTEIN TC10. Length = 213		antigenic surface determinant OA3 [Homo sapiens] >pir[A48997 A48997 tumor surface antigen OA3-323 - human >sp Q08722 CD47_HUMAN LEUKOCYTE SURFACE ANTIGEN CD47 PRECURSOR (ANTIGENIC SURFACE DETERMINANT PROTEIN OA3) (INTEGRIN ASSOCIATED PROTEIN) (IAP) (MER6)	insulin-like growth factor-binding protein [Homo sapiens] >gb AAA52706.1  growth factor-binding protein-3 [Homo sapiens] >emb CAA46087.1  insulin-like growth factor binding protein 3 [Homo sapiens] >pir A36578 IOHU3 insulin-like growth factor-binding prot	(AJ002190) dihydroxyacetone phosphate acyltransferase [Homo sapiens] >gb AAC24505.1  (AF043937) peroxisomal acyl-CoA:dihydroxyacetonephosphate acyltransferase [Homo sapiens] >sp O15228 DAPT_HUMAN DIHYDROXYACETONE PHOSPHATE ACYLTRANSFERASE (EC 2.3.1.42) (D	protein tyrosine phosphatase [Homo sapiens] >gb AAA66496.1  protein phosphatase [Homo sapiens] >sp Q16667 CDN3_HUMAN CYCLIN- DEPENDENT KINASE INHIBITOR 3 (EC 3.1.3.48) (EC 3.1.3.16) (CDK2- ASSOCIATED DUAL SPECIFICITY PHOSPHATASE) (KINASE ASSOCIATED PHOSPHA
877361	877363	877370	877373	877375	877377
1891	1892	1893	1894	1895	1896

pBluescript SK-	Uni-ZAP XR	HELBN30 Uni-ZAP XR	HHFMH12 Uni-ZAP XR	HBXAC19 ZAP Express	pSport1
HILBZ32	HAPOR25	HELBN30	HHFMH12	HBXAC19	HWLNV37
66	77	84	72	100	100
66	77	84	24	100	100
605	1782	1035	2161	81	096
27	-	157	2	_	499
6174	6175	6176	6177	6178	6179
gb AAA60222.	emb CAA5208	gb AAA35537. 1	dbj BAA91631 .1	gb AAB05170.	gb AAF03505. 1 AC0049
protein tyrosine phosphatase [Homo sapiens] >gb AAA66496.1  protein phosphatase [Homo sapiens] >sp Q16667 CDN3_HUMAN CYCLIN- DEPENDENT KINASE INHIBITOR 3 (EC 3.1.3.48) (EC 3.1.3.16) (CDK2- ASSOCIATED DUAL SPECIFICITY PHOSPHATASE) (KINASE ASSOCIATED PHOSPHA	microtubule associated protein [Homo sapiens] emb >pir 137356 137356 epithelial microtubule-associated [6.1] protein, 115K - human >sp Q14244 Q14244 MICROTUBULE ASSOCIATED PROTEIN (DJ406A7.2.1) (MICROTUBLE ASSOCIATED PROTEIN E-MAP-115). >emb CAB37984.1  (AL023284)	nuclear autoantigen [Homo sapiens] >pir A37244 A37244 nuclear autoantigen Sp-100 - human Length = 480	(AK001332) unnamed protein product [Homo sapiens] Length = 682	receptor tyrosine kinases AC51203.1  putative EPH- igand LERK-8 [Homo sapiens] JMAN EPHRIN-B3 KELATED RECEPTOR LIGAND 8) (LERK-8) (EPH-	(AC004922) similar to G10 protein; similar to AAC14190 (PID:g3064070) [Homo sapiens] >sp AAF03505 AAF03505 WUGSC:H_DJ0900K19.2 protein. Length = 144
877378	877380	877384	877387	877388	877390
1897	1898	1899	1900	1901	1902

HWHQHI7 pCMVSport 3.0	HDPFP36 pCMVSport 3.0	HCFMY07 pSport1	HSYBP46 pCMVSport 3.0	HCRQK59 pSport1	HWLXK44 pSport1	HE8DZ94 Uni-ZAP XR	HTELO87 Uni-ZAP XR	HWLQL72 pSport1
Н 96		94	1000	-	100 H	91		1
96		93	100		100	91		
2530	804	2498	1013	313	314	1657	662	553
1205	340	1296	E.	2	93	1256	ε.	245
6180	6181	6182	6183	6184	6185	6186	6187	6188
gb AAA51783. 1		gb AAB02814. 1	emb CAA7500		gb AAC51782. 1	gb AAD34114. 1 AF1518		
argininosuccinate synthetase [Homo sapiens] >emb CAA25771.1  argininosuccinate synthetase (aa 1-412) [Homo sapiens] >pir A01195 AJHURS argininosuccinate synthase (EC 6.3.4.5) - human >sp P00966 ASSY_HUMAN ARGININOSUCCINATE SYNTHASE (EC 6.3.4.5) (CITRULLIN		Rho-associated, coiled-coil containing protein kinase p160ROCK [Homo sapiens] >pir S69211 S69211 serine/threonine-specific protein kinase (EC 2.7.1), Rho-associated - human >sp Q13464 Q13464 RHO-ASSOCIATED, COILED-COIL CONTAINING PROTEIN KINASE P160ROCK	procollagen alpha 2(V) [Homo sapiens] >pir A31427 CGHU2V collagen alpha 2(V) chain precursor - human >sp P05997 CA25_HUMAN COLLAGEN ALPHA 2(V) CHAIN PRECURSOR. >sp CAA75002 CAA75002 Procollagen alpha 2(V). >emb CAA28454.1  pro- alpha (V)collagen (AA 1099)		multispanning membrane protein [Homo sapiens] >sp O15321 O15321 MULTISPANNING MEMBRANE PROTEIN. Length = 606	(AF151877) CGI-119 protein [Homo sapiens] >sp Q9Y3C2 Q9Y3C2 CGI-119 PROTEIN. >gb AAF14868.1 AF113127_1 (AF113127) S1R protein [Homo sapiens] {SUB 21-258} Length = 258		
877393	877396	877406	877408	877411	877437	877630	877881	878199
1903	1904	1905	1906	1907	1908	1909	1910	1911

1912	878207	(AK001523) unnamed protein product [Homo sapiens] Length = 165	dbj BAA91739 .1	6189	572	1069	68	68	HBJJL05	Uni-ZAP XR
1913	878238	ister trithorax protein is] >pir T34384 T34384 26A5.7 - Caenorhabditis 22795 HYPOTHETICAL 27.0	gb AAC77512. 1	6190	3	272	09	73	HE2HC14	Uni-ZAP XR
1914	878274	(AF203978) MAX-like bHLHZIP protein [Homo sapiens] >sp AAF14638 AAF14638 MAX-like bHLHZIP protein. Length = 244	gb AAF14638. 1 AF2039	6191	2	487	100	100	HDTHI51	pCMVSport 2.0
1915	878374	(AB033010) KIAA1184 protein [Homo sapiens] >sp BAA86498 BAA86498 KIAA1184 protein (fragment). Length = 380	dbj BAA86498 .1	6192	2	1051	68	68	HRGDE77	HRGDE77 Uni-ZAP XR
1916	878403	3-hydroxy-3-methylglutaryl coenzyme A synthase [Homo sapiens] >pir[845497]845497 hydroxymethylglutaryl-CoA synthase (EC 4.1.3.5), cytosolic, adrenal isoform - human >sp[Q01581 HMCS_HUMAN HYDROXYMETHYLGLUTARYL-COA SYNTHASE, CYTOPLASMIC (EC 4.1.3.5) (HMG-CO	gb AAA62411. 1	6193	283	882	91	93	ннғнкѕз	Uni-ZAP XR
1917	878433	(AF096895) chemokine-like factor 1 [Homo sapiens] gb AAF06722. >sp AAF06722 AAF06722 Chemokine-like factor 1. 1 AF0968 Length = 99	gb AAF06722; 1 AF0968	6194	105	443	100	100	HTPAY82	Uni-ZAP XR
1918	878436	(AK001682) unnamed protein product [Homo sapiens] Length = 242	dbj BAA91833 .1	6195	82	1524	66	66	НМИВQ39	pCMVSport 3.0
1919	878560	(AF108139) radical fringe [Homo sapiens] >sp Q9Y644 Q9Y644 RADICAL FRINGE (FRAGMENT). Length = 191	gb AAD34321. 1 AF1081	6196	331	2	86	100	HCEYN60	Uni-ZAP XR
1920	878800	ORF_f418 [Escherichia coli] >gb AAD13442.1  (AE000464) orf, hypothetical protein [Escherichia coli] >pir S40824 S40824 hypothetical 48K protein (glnA-fdhE intergenic region) - Escherichia coli >sp P32140 YIHS_ECOLI HYPOTHETICAL 47.4 KD PROTEIN IN GLNA-RBN	gb AAB03013. 1	6197	223	2	100	100	HWHGF46	pCMVSport 3.0

1921	878909	KIAA0182 [Homo sapiens] >sp[Q14687]Y182_HUMAN HYPOTHETICAL PROTEIN KIAA0182 (FRAGMENT). Length = 1157	dbj BAA11499 .1	6198	699	1421	84	85	HPMSF50	pBluescript
1922	878917	KIAA0069 [Homo sapiens] >sp[Q15041 Y069_HUMAN HYPOTHETICAL PROTEIN KIAA0069 (HA1508) (FRAGMENT). Length = 226	dbj BAA06683 .1	6199	2	346	77	77	HTWEA61	pSport1
1923	878931			9700	318	909	L.		HILBF77	pBluescript SK-
1924	879009	(AF035606) calcium binding protein [Homo sapiens] gb AAC27697. >gb AAF14336.1 U58773_1 calcium binding protein 1  [Homo sapiens] >sp O75340 O75340 CALCIUM BINDING PROTEIN. >sp AAF14336 AAF14336 Calcium binding protein. Length = 191	gb AAC27697. 1	6201	618	1040	100	100	нтенх05	HTEHX05 Uni-ZAP XR
1925	879234			6202	211	903			HPHAA47	Uni-ZAP XR
1926	879386	(AF161516) HSPC167 [Homo sapiens] >sp AAF29131 AAF29131 HSPC167. Length = 586	gb AAF29131. 1 AF1615	6203	419	991	100	100	HHFJJ61	Uni-ZAP XR
1927	879484	(AF053651) cellular apoptosis susceptibility protein [Homo sapiens] >sp 075432 075432 CELLULAR APOPTOSIS SUSCEPTIBILITY PROTEIN. Length = 971	gb AAC35297. 1	6204	885	2108	100	100	H2CAA49	pBluescript SK-
1928	879595			6205	3	419			HCRNW08	pSport1
1929	879661	(AF151079) HSPC245 [Homo sapiens] Length = 124 gb AAF36165.	gb AAF36165. 1 AF1510	6206	158	982	100	100	HNTDJ29	pCMVSport 3.0
1930	988628			6207	502	762			HCRNM29	pSport1
1931	880071			6208	292	921			HTPAM76	Uni-ZAP XR
1932	880074	(AF112214) ribosomal protein L13 [Homo sapiens] >sp AAF17202 AAF17202 Ribosomal protein L13. Length = 172	gb AAF17202. 1 AF1122	6209	290	829	66	100	нснов95	pSport1
1933	880418			6210	1082	1339			HLSAA96	pSport1

pCMVSport	HE8QG48 Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	pCMVSport 2.0	HOSML44 Uni-ZAP XR	Uni-ZAP XR	Uni-ZAP XR	pBluescript
HBBMA61	HE8QG48	HHENW13	HE8SB64	HKAEN78	HOSML44	HTEEZ62	НОААН52	HSDXB50
	100	88	77			96	97	92
	100	88	74			96	97	92
255	998	618	1751	922	527	1565	1277	654
130	09		3	422	339	m	8	265
6211	6212	6213	6214	6215	6216	6217	6218	6219
	gb AAD26810. 1 AF1192	dbj BAA25263 .1	dbj BAA18909 .1			dbj BAA74873 .1	dbj BAA19780  .1	emb CAB7536 7.1
	(AF119297) neuroendocrine-specific protein-like protein 1 [Homo sapiens] >gb AAC99319.1  (AF059524) reticulon gene family protein [Homo sapiens] >gb AAD20951.1  (AF059529) reticulon gene family protein [Homo sapiens] >sp O95197 O95197 RETICULON PROTEIN. L	(AB001740) p27 [Homo sapiens] db   >sp O60232 AA27_HUMAN AUTOANTIGEN P271  Length = 199	ns] >sp Q14089 Q14089  CAL 40.0 KD PROTEIN  C. Length = 364			(AB020657) KIAA0850 protein [Homo sapiens] >emb CAB72329.1  (AL078644) bG279B7.1.1 (NS1-binding protein (KIAA0850, BTB/POZ domain and Kelch motifs containing protein)) [Homo sapiens] >sp Q9Y6Y0 Q9Y6Y0 KIAA0850 PROTEIN. Length = 642	Similar to a C.elegans protein in cosmid C14H10 [Homo sapiens] >sp O00236 O00236 KIAA0251 (FRAGMENT). Length = 820	(AL050318) dJ977B1.3.1 (novel protein similar to putative RAB5-interacting protein (isoform 1)) [Homo sapiens] >gb AAF17201.1 AF112213_1 (AF112213) putative Rab5-interacting protein [Homo sapiens] >sp AAF17201 AAF17201 Putative Rab5-interacting protein. L
880578	880649	880694	880747	880927	880994	881052	881074	881104
1934	1935	1936	1937	1938	1939	1940	1941	1942

HFKMJ24 Uni-ZAP XR	pSport1	pSport1	Uni-ZAP XR	HMEKW44 Lambda ZAP II	Uni-ZAP XR	pSport1	pSport1	pSport1	Other	pSport1
HFKMJ24	необси	HWMBI22	HETDL42	HMEKW44	HCEDM42	HCRNZ31	HWMBU89	HUFBY15	HIBCE91	HWLKF77
100	95	92		74	08				71	85
100 100	95	92		72	80				71	82
576	2348	1036	1477	327	992	489	652	468	569	917
127	$\epsilon$	2	1079	1	110	1	86	295	3	3
6220	6221	6222	6223	6224	6225	6226	6227	6228	6229	6230
emb CAB7536	gb AAF03515. 1 AC0048	gb AAC50897. 1		emb CAB0666 4.1	gb AAF36109. 1 AF1510				gb AAC31671. 1	emb CAB6510 5.1
(AL050318) dJ977B1.3.1 (novel protein similar to putative RAB5-interacting protein (isoform 1)) [Homo sapiens] >gb AAF17201.1 AF112213_1 (AF112213) putative Rab5-interacting protein [Homo sapiens] >sp AAF17201 AAF17201 Putative Rab5-interacting protein. L	(AC004882) similar to CAA16821 (PID:g3255952) [Homo sapiens] >sp AAF03515 AAF03515 WUGSC:H_DJ076B20.5 protein (fragment). Length = 620	cell growth regulator CGR19 [Homo sapiens] >sp Q99675 Q99675 CELL GROWTH REGULATOR CGR19. Length = 332		hypothetical protein [Bos taurus] >sp O18975 O18975 HYPOTHETICAL 16.6 KD PROTEIN (FRAGMENT). Length = 145	(AF151023) HSPC189 [Homo sapiens] Length = 222 gb AAF36109.				(AC003007) gene product (partial) [Homo sapiens] >sp 075201 075201 HYPOTHETICAL 64.6 KD PROTEIN (FRAGMENT). Length = 580	(AJ245719) brk kinase substrate [Homo sapiens] >sp CAB65105 CAB65105 Brk kinase substrate. Length = 403
881105	881219	881221	882330	882715	882729	882762	883172	883201	883254	883371
1943	1944	1945	1946	1947	1948	1949	1950	1951	1952	1953

pCMVSport 2.0	pCMVSport 2.0	pSport1	Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	pSport1	Uni-ZAP XR	pSport1	pSport1	pCMVSport 3.0	pSportl
HOGCA75 pCMVSport 2.0	HOGCJ47	HWLUT61	HLTBA42	нненв82	HE2PR08	HMKAN71	HSIFV30	HNTSY52	HCROM43	HLWCF60	HWLKD85
77				93	72	100	96	06	75	100	71
77				93	46	99	96	81	73	100	89
898	1129	157	303	741	956	1385	2776	1171	775	683	436
239	998	65	115	208	432	3	2	239	2	237	59
6231	6232	6233	6234	6235	6236	6237	6238	6239	6240	6241	6242
dbj BAA22984				gb AAD34119.  1 AF1518	dbj BAA91309 .1	dbj BAA91938 .1	gb AAD25487. 1 AF1270	dbj BAA85045 .1	gb AAF06800. 1 AF1969	gb AAD34145. 1 AF1519	dbj BAA88307 .1
(AB000712) CPE-receptor [Homo sapiens] >sp 014493 CLD4_HUMAN CLAUDIN-4 (CLOSTRIDIUM PERFRINGENS ENTEROTOXIN RECEPTOR) (CPE- RECEPTOR) (CPE-R). Length = 209				protein [Homo sapiens] CGI-124 PROTEIN (EC	(AK000654) unnamed protein product [Homo sapiens] Length = 248	(AK001845) unnamed protein product [Homo sapiens] Length = 612	ctivated chloride channel ns] 55487 Calcium-activated in 1. Length = 914	Homo ein kinase	(AF196972) JM24 protein [Homo sapiens]       gb AAF06800.         >sp AAF06800 AAF06800 JM24 protein (fragment).       I: AF1969         Length = 476	(AF151908) CGI-150 protein [Homo sapiens] >sp Q9Y3E8 Q9Y3E8 CGI-150 PROTEIN. Length = 504	(AB028859) hDj9 [Homo sapiens] dt >emb CAB65118.1  (AJ250137) ERj3 protein [Homo .1 sapiens] >sp CAB65118 CAB65118 ERj3 protein precursor. >sp BAA88307 BAA88307 HDj9. Length = 358
883753	883799	883945	883971	884038	884095	884161	884168	884215	884379	884529	884719
1954	1955	1956	1957	1958	1959	1960	1961	1962	1963	1964	1965

1966	885350	(AB011532) MEGF6 [Rattus norvegicus] > pir T13954 T13954 MEGF6 protein - rat > sp O88281 O88281 MEGF6. Length = 1574	dbj BAA32462 .1	6243	430	1581	45	95	HCRMX54	pSport1
1967	885476			6244	432	806		<u> </u>	HTPHK88	Uni-ZAP XR
1968	885484	ORF4 [Rattus norvegicus] >pir S21348 S21348 probable pol polyprotein-related protein 4 - rat >sp Q63306 Q63306 LONG INTERSPERSED REPETITIVE DNA CONTAINING 7 ORF"S. Length = 275	emb CAA3764 7.1	6245	396	476	42	73	нсоврзя	HCQBD35 Lambda ZAP
1969	885511			6246	178	408			HLQF167	Lambda ZAP II
1970	886331	(AF026124) schwannoma-associated protein [Musmusculus] >sp 035405 035405 SCHWANNOMA-ASSOCIATED PROTEIN. Length = 488	gb AAC73069.	6247	3	755	37	57	HAJBV26	pCMVSport 3.0
1971	886505	(AF161410) HSPC292 [Homo sapiens] >sp AAF28970 AAF28970 HSPC292 (fragment). Length = 164	gb AAF28970. 1 AF1614	6248	565	975	100	100	HBJJF90	Uni-ZAP XR
1972	886527			6549	1	375			HWLFB44	pSport1
1973	886788	(AK001350) unnamed protein product [Homo sapiens] Length = 326	dbj BAA91642 .1	6250	96	965	86	66	HCE4U96	Uni-ZAP XR
1974	886914	A33 antigen precursor [Homo sapiens] >sp Q99795 A33 HUMAN CELL SURFACE A33 ANTIGEN PRECURSOR. Length = 319	gb AAC50957. 1	6251	3	443	100	100	HWLEL48	pSport1
1975	887098	(AF161453) HSPC335 [Homo sapiens] >sp AAF29013 AAF29013 HSPC335 (fragment). Length = 159	gb AAF29013. 1 AF1614	6252	3	539	68	06	HTGBT14	Uni-ZAP XR
1976	887114	(AF067797) aquaporin 8 [Homo sapiens] >sp AAF19050 AAF19050 Aquaporin 8. Length = 261	gb AAF19050. 1	6253	297	1160	83	83	HKLRB09	pBluescript
1977	887155			6254	3	497			H2LAS29	pBluescript SK-
1978	887172	(AK000700) unnamed protein product [Homo sapiens] Length = 370	dbj BAA91327 .1	6255	505	1071	06	92	нмекн10	Lambda ZAP II

pSport1	pBluescript	pBluescript	pSport1	pCMVSport 3.0	Uni-ZAP XR	pBluescript	pCMVSport 1
HWLWR39	HADME31	HFVJL45	HWLFE56	HSWBP93	HSLJF91	HKLSC61	HLJEA63
96		66	86		68		100
95		66	86		98		100
1205	793	1134	281	475	464	757	1059
195	527	25	က	218	180	377	319
6256	6257	6258	6259	6260	6261	6262	6263
emb CAA0082 9.1		sp[Q15166]PO	emb CAB5548 9.1		emb CAA5714 3.1		emb CAA0962  6.1
urokinase [synthetic construct] >emb CAA00996.1  human u-PA cDNA insert [synthetic construct] >emb CAA01390.1  uPA [Homo sapiens] >emb CAA01559.1  pro-Urokinase [Homo sapiens] >emb CAA02215.1  u-PA [Homo sapiens] >dbj BAA00175.1  pro-urokinase precursor [		SERUM PARAOXONASE/ARYLESTERASE 3 (EC 3.1.1.2) (EC 3.1.8.1) (PON 3) (SERUM ARYLDIAKYLPHOSPHATASE 3) (A-ESTERASE 3) (AROMATIC ESTERASE 3). >gb AAC62430.1  (AC005021) serum paraoxonase/arylesterase [Homo sapiens] {SUB 1-122} Length = 354	(AJ009936) nuclear hormone receptor PRR1-A [Homo sapiens] >sp CAB55489 CAB55489 Nuclear hormone receptor PRR1-A. >emb CAB55491.1  (AJ009936) nuclear hormone receptor PRR1-B [Homo sapiens] {SUB 56-434} Length = 434		unnamed protein product [Sus scrofa] >pir S52130 S52130 vascular endothelial growth factor - pig >sp P49151 VEGF_PIG VASCULAR ENDOTHELIAL GROWTH FACTOR PRECURSOR (VEGF) (VASCULAR PERMEABILITY FACTOR) (VPF). Length = 190		(AJ011497) Claudin-7 [Homo sapiens] >sp O95471 CLD7_HUMAN CLAUDIN-7. Length = 211
887192	887280	887399	887421	887475	887535	887803	887857
1979	1980	1981	1982	1983	1984	1985	1986

	100 HCQCF10 Lambda ZAP	86 HAIBW90 Uni-ZAP XR	97 H2CBE03 pBluescript SK-	64 HE9Q119 Uni-ZAP XR	91 HJACE25 pBluescript SK-	80 HMWIR85 Uni-ZAP XR
59	100	98	76	62	91	08
519	199	403	604	1066	399	1907
82	7	7	2	2		510
6264	6265	6266	6267	6268	6569	6270
emb CAA6064 5.1	gb AAD20035.	.1	dbj BAA11492 .1	gb AAB58505. 1	gb AAC64044. 1	dbj BAA37094 .1
2.19 [Homo sapiens] >emb CAA39090.1  2-19 protein [Homo sapiens] >gb AAA92652.1  2_19 [Homo sapiens] >pir [37095 [37095 gene 2.19 protein - human >sp P98173 219 HUMAN 2-19 PROTEIN PRECURSOR. Length = 230	(AF131758) [Homo sapiens] >sp O95881 O95881. Length = 172	(AB006077) deleted in oral cancer 1 (doc-1, alias DORC1) [Homo sapiens] >gb AAC77831.1] (AF006484) putative oral tumor suppressor protein [Homo sapiens] >sp O14519 DOC1_HUMAN PUTATIVE ORAL CANCER SUPPRESSOR (DELETED IN ORAL CANCER-1). Length = 115	similar to protein kinase of X.laevis, has putative transmembrane domain incentral region [Homo sapiens] >sp Q14680 Q14680 KIAA0175 PROTEIN. Length = 651	fatty acid amide hydrolase [Homo sapiens] >gb AAD13768.1  (AF098019) fatty acid amide hydrolase [Homo sapiens] >sp O00519 FAAH_HUMAN FATTY ACID AMIDE HYDROLASE (EC 3.1) (OLEAMIDE HYDROLASE). Length = 579	(AF073771) RNA polymerase II termination factor [Homo sapiens] >sp 075921 075921 RNA POLYMERASE II TERMINATION FACTOR. Length = 1162	(AB013357) 49 kDa zinc finger protein [Mus musculus] >pir JE0367 JE0367 zinc finger protein - mouse >sp Q9Z326 Q9Z326 49 KDA ZINC FINGER PROTEIN. Length = 460
887892	887936	984388	888041	888051	888063	888153
1987	1988	6861	1990	1991	1992	1993

pSport1	Uni-ZAP XR	pCMVSport 3.0	Uni-ZAP XR	Uni-ZAP XR	pCMVSport 3.0	pBluescript SK-
HCRPV38	HSRBB92	HSYEA10	HE2CC22	HOUAC22	HHECU01	H2LAP34
100	93	100		62		89
100	93	100		48		40
334	2011	1489	286	645	964	371
2	2	2	2	241	488	111
6271	6272	6273	6274	6275	6276	6277
gb AAB63956. 1	emb CAB6770	emb CAA3869		emb CAA1593		gb AAA72205.
glutathione S-transferase theta 2 [Homo sapiens] >pir A56847 A56847 glutathione transferase (EC 2.5.1.18) theta-2 - human >sp P30712 GTT2_HUMAN GLUTATHIONE S-TRANSFERASE THETA 2 (EC 2.5.1.18) (GST CLASS-THETA). {SUB 2-244} Length = 244	(AJ271408) Fas-associated factor, FAFI [Homo eml sapiens] >gb AAD27713.1 AF132938_1 (AF132938) 5.1  CGI-03 protein [Homo sapiens] >sp Q9Y2Z3 Q9Y2Z3 CGI-03 PROTEIN. >sp CAB67705 CAB67705 Fas-associated factor, FAFI. >emb CAB63755.1  (AL133631) hypothetical protein	type I interstitial collagenase [Homo sapiens] >gb AAB36941.1  collagenase [Homo sapiens] >pir A37308 KCHUI interstitial collagenase (EC 3.4.24.7) precursor - human >sp P03956 COG1_HUMAN INTERSTITIAL COLLAGENASE PRECURSOR (EC 3.4.24.7) (MATRIX METALLOPROT		(AL021106) /prediction=(method:"genscan", version:"1.0", score:"113.71"); /prediction=(method:"genefinder", version:"084"); /match=(desc:"LD30851.5prime LD Drosophila melanogaster embryo pOT2 Drosophila melanogaster cDNA clone LD30851 5prime>>		calcium-modulated protein S100-beta [synthetic construct] >pir A91254 BCBOIB S-100 protein beta chain - bovine {SUB 2-92} Length = 92
888254	888402	888523	888673	888708	888720	888783
1994	1995	1996	1997	1998	6661	2000

2001	888950	growth factor [Mus musculus] >pir A46607 A46607 growth/differentiation factor GDF-3 precursor -mouse >sp Q07104 GDF3_MOUSE GROWTH/DIFFERENTIATION FACTOR 3 PRECURSOR (GDF-3) (VG-1-RELATED PROTEIN 2). Length = 366	gb AAA53034. 1	6278	<b>E</b>	1151	70	08	HNTAR08	pCMVSport 3.0
2002	889136			6279	311	520			99НМТМН	pSport1
2003	889263	beta-galactosidase [Expression vector pBSII- LUCINT] Length = 69	gb AAB53629.	6280	1924	2259	94	86	HWLCJ12	pSport1
2004	889299			6281	1329	1520			HNGEF72	Uni-ZAP XR
2005	889300	(AB011145) KIAA0573 protein [Homo sapiens] >sp O60319 O60319 KIAA0573 PROTEIN (FRAGMENT). Length = 451	dbj BAA25499 .1	6282	3	1409	97	97	HKAEB46	pCMVSport 2.0
2006	889323			6283	481	966			HNHON23	Uni-ZAP XR
2007	889368	SThM [Homo sapiens] >sp[Q12971 Q12971 SIALYLTRANSFERASE STHM. Length = 374	gb AAA52228. 1	6284	3595	3176	86	86	HSKES11	Uni-ZAP XR
2008	889467	(AL096745) hypothetical protein [Homo sapiens] >emb CAB62532.1  (AL096745) hypothetical protein [Homo sapiens] >pir T12548 T12548 hypothetical protein DKFZp586D1022.1 - human >sp CAB62532 CAB62532 Hypothetical 40.6 kd protein. Length = 366	emb CAB6253 2.1	6285	r.	413	86	66	<b>НСЕТР05</b>	pBluescript
2009	889494	(AF065389) tetraspan NET-4 [Homo sapiens] >gb AAF28869.1 AF121344_1 (AF121344) tetraspanin Tspan-5 [Mus musculus] >sp O60746 O60746 TETRASPAN NET-4. >sp AAF28869 AAF28869 Tetraspanin Tspan-5. Length = 268	gb AAC17120. 1	6286	2	361	74	78	нрнел	pCMVSport 2.0
2010	889700			6287	1	459			HCHAC08	pSport1
2011	889782	aldehyde dehydrogenase [Homo sapiens] Length = 517	gb AAA51693. 1	6288	140	463	100	100	HACBT96	Uni-ZAP XR

HTLEN01 Uni-ZAP XR	pSport1	HSLJW05 Uni-ZAP XR	Uni-ZAP XR	Lambda ZAP II	HOSOR86 Uni-ZAP XR	HE9RV77 Uni-ZAP XR	HPRAJ70 Uni-ZAP XR
HTLEN01	HCROA43	HSLJW05	HTPGK74	HHGAB64	HOSOR86	HE9RV77	HPRAJ70
92	47		82	69		100	97
85	34		88	99		100	93
1025	705	446	1308	256	1632	854	2420
3	1	282	46	17	1399	E.	1317
6289	96790	6291	6292	6293	6294	6295	6296
dbj BAA86538 .1	gb AAC69835. 1		6.1	emb CAB4015		emb CAB5568	gb AAD12761. 1
224 protein [Homo sapiens] 86538 KIAA1224 protein 635	(AF076612) chordin [Homo sapiens] >sp O95254 O95254 CHORDIN (FRAGMENT). Length = 801		membrane cofactor preprotein (AA -34 to 350) [Homo sapiens] >pir S01896 S01896 membrane cofactor protein precursor - human >gb AAD13968.1 S65879_1 complement system membrane cofactor protein CD46 [Homo sapiens] {SUB 1-34} Length = 384	(AL031295) dJ886K2.3(GALE (UDP-galactose-4-epimerase)) [Homo sapiens] >sp CAB40159 CAB40159 DJ886K2.3(GALE (UDP-galactose-4-epimerase)). Length = 348		(AL035608) dJ479J7.2 (transmembrane 4 superfamily member 6) [Homo sapiens] >gb AAC64257.1  (AF043906) T245 protein [Homo sapiens] >gb AAC69710.1  (AF053453) tetraspan TM4SF [Homo sapiens] >gb AAD00560.1  A15 homolog [Homo sapiens] >gb AAF08365.1 AF133426_	(AF079864) putative G-protein coupled receptor RA1c [Rattus norvegicus] >spl088628 088628 PUTATIVE G-PROTEIN COUPLED RECEPTOR RA1C. Length = 320
889954	889962	889994	999068	869068	890753	890763	890776
2012	2013	2014	2015	2016	2017	2018	2019

2020	890801	(AB035207) Tob2 [Homo sapiens] >emb CAB62938.1  (AL008582) bK223H9.1 (TOB4 (BTG1 family protein)) [Homo sapiens] >sp BAA87042 BAA87042 Tob2. >sp CAB62938 CAB62938 BK223H9.1 (TOB4 (BTG1 family protein)). Length = 344	dbj BAA87042 .1	6297	501	1505	9	H 59	HBODK52	pSport1
2021	890820	(AF009702) GABA-A receptor pi subunit [Homo sapiens] >gb AAC51357.1  GABA-A receptor pi subunit [Homo sapiens] >sp 000591 GAAP_HUMAN GAMMA-AMINOBUTYRIC-ACID RECEPTOR PI SUBUNIT PRECURSOR (GABA(A) RECEPTOR). Length = 440	gb AAC24194. 1	6298	134	1516	95	95 H	HARNK52	pCMVSport 3.0
2022	890863	(AK000207) unnamed protein product [Homo sapiens] Length = 478	dbj BAA91009 .1	6299	2	1210	45	64 F	TLHU22	HTLHU22 Uni-ZAP XR
2023	890945	DNA mismatch repair protein homolog [Homo sapiens] >gb AAC50285.1  hMLH1 [Homo sapiens] >pir S43085 S43085 DNA mismatch repair protein MLH1 - human >sp P40692 MLH1 HUMAN MUTL PROTEIN HOMOLOG 1 (DNA MISMATCH REPAIR PROTEIN MLH1). Length = 756	gb AAA82079. 1	6300	3	2327	96	Н 96	HWMBB29	pSport1
2024	891125	pancreatic peptidylglycine alpha-amidating monooxygenase, PAM=secretory isoform {clone PAM-15} [human, islet cell tumor cell line QGP-1, Peptide Partial, 905 aa] [Homo sapiens] >sp Q16253 Q16253 PANCREATIC PEPTIDYLGLYCINE ALPHA-AMIDATING MONOOXYGENASE (FR	gb AAB32776. 1	6301	2	478	86	H 86	HWLND63	pSport1
2025	891264	(AK001537) unnamed protein product [Homo sapiens] Length = 129	dbj BAA91746 .1	6302	_	633	001	100 H	нскод71	pSport1
2026	891305	piens] JMAN HYPOTHETICAL (HA6725). Length = 247	dbj BAA13397 .1	6303	227	2338	55	75 1	HBINP81	pCMVSport 3.0
2027	891896	beta-galactosidase alpha peptide [Cloning vector pSport2] Length = 114	gb AAA67217. 1	6304	2	226	69	80	HDLAG89	pCMVSport 2.0

2028	892113	conserved hypothetical protein MTH68 - Methanobacterium thermoautotrophicum (strain Delta H) Length = 228	pir E69190 E6 9190	6305	844	1557	23	42	HE8FL95	HE8FL95 Uni-ZAP XR
2029	892177	cal protein [Homo sapiens]  L137599) hypothetical  J-sp CAB70835 CAB70835  protein (fragment). Length =	emb CAB7083 5.1	6306	33	1067	57	71	HHFGI59	Uni-ZAP XR
2030	892291	cathepsin D [Homo sapiens] >emb CAA28955.1  precursor polypeptide (AA -20 to 392) [Homo sapiens] >gb AAB59529.1  preprocathepsin D [Homo sapiens] >pir A25771 KHHUD cathepsin D (EC 3.4.23.5) precursor - human >sp P07339 CATD_HUMAN CATHEPSIN D PRECURSOR (EC	gb AAA51922.	6307	21	1232	96	96	HOFMT75	pCMVSport 2.0
2031	892367	!!!! ALU SUBFAMILY SC WARNING ENTRY !!!! sp P39192 AL Length = 585	sp P39192 AL US_HUMAN	6308	668	1018	78	48	HWLEQ37	pSport1
2032	892558			6309	770	926			HWLDZ74	pSport1
2033	892563	MAL protein [Homo sapiens] >emb CAA53809.1  MAL [Homo sapiens] >emb CAA54100.1  MAL-a [Homo sapiens] >pir A29472 A29472 T-cell surface glycoprotein MAL, splice form a - human >sp P21145 MAL_HUMAN MYELIN AND LYMPHOCYTE PROTEIN (T-LYMPHOCYTE MATURATION-ASSO	gb AAA36196.	6310	_	618	39	50	HPJEB77	Uni-ZAP XR
2034	892820			6311	371	553			HNTST71	pSport1
2035	893223	(AF216312) type II membrane serine protease [Homo sapiens] Length = 423	gb AAF31436. 1 AF2163	6312	2	508	66	66	нсороэ2	HCQDQ92 Lambda ZAP
2036	893457			6313	88	345			HWLCU24	pSportl

HSDJY15 Uni-ZAP XR	pBluescript SK-	6 pCMVSport 2.0	3 pBluescript	HSVCD79 Uni-ZAP XR	Uni-ZAP XR	HTTKV46 Uni-ZAP XR
HSDJY15	HSAAR81	HNDAD16	HCNSE58	HSVCD79	HSIFA27	HTTKV46
72		78	78	100		87
71		78	78	86		87
829	388	436	316	257	310	855
512	98	41	2	51	23	
6314	6315	6316	6317	6318	6319	6320
emb CAA2403 8.1		gb AAA59981. 1	gb AAA59981. 1	gb AAC63910. 1		pir B42856 B4 2856
cytochrome B [Homo sapiens] >pir A00151 CBHU ubiquinolcytochrome-c reductase (EC 1.10.2.2) cytochrome b - human mitochondrion >sp P00156 CYB_HUMAN_CYTOCHROME_B. >gb AAB58955.1  cytochrome b [Homo sapiens] {SUB 1-378} >gb AAA31851.1  cytochrome b [Homo s		secretory protein [Homo sapiens] >gb AA83628.1  intestinal trefoil factor [Homo sapiens] >pir A48284 A48284 intestinal trefoil factor 3 precursor - human >sp Q07654 ITF_HUMAN INTESTINAL TREFOIL FACTOR PRECURSOR (HP1.B). Length = 80	secretory protein [Homo sapiens] >gb AAA83628.1  intestinal trefoil factor [Homo sapiens] >pir A48284 A48284 intestinal trefoil factor 3 precursor - human >sp Q07654 ITF_HUMAN INTESTINAL TREFOIL FACTOR PRECURSOR (HP1.B). Length = 80	(AF023259) RNA-binding protein [Homo sapiens] >gb AAC63910.1  (AF023259) RNA-binding protein [Homo sapiens] >sp O75876 O75876 RNA-BINDING PROTEIN. Length = 411		ubiquitin carrier protein E2 - human >gb AAA58446.1  ubiquitin carrier protein [Homo sapiens] {SUB 23-247} Length = 247
893827	893842	893866	893867	894012	894051	894121
2037	2038	2039	2040	2041	2042	2043

a ZAP I	script <-	ort1	ortl	ortl	la ZAP I	pBluescript	uescript SK-	la ZAP I	da ZAP II
Lambd	pBluescript SK-	pSport1	pSport1	pSport1	Lambda II	pBlue	pBluescript SK-	Lambd I	Lambd
HHGCE29 Lambda ZAP	HCYBE73	HWLVS05	HCRMV27	HCROI22	HCQAF06 Lambda ZAP	HKCSA83	HSBAI04	HCQCD80 Lambda ZAP	HCQCF52 Lambda ZAP
100	72	98				100			66
100	42	98				100			66
122	355	1268	411	620	102	576	3	570	684
ĸ	2	54	70	381	-	373	77	253	244
6321	6322	6323	6324	6325	6326	6327	6328	6329	6330
dbj BAA01133   -1	dbj BAA89784 .1	emb CAB5926 6.1				dbj BAA16267			dbj BAA28861
homeodomain protein [Gallus gallus] >pir B37914 B37914 homeotic protein Chox-4e - chicken (fragment) {SUB 7-99} >pir S14938 S14938 homeotic protein Hox D11 - human (fragment) {SUB 27-92} Length = 99	(AB037134) IRE homolog 1 [Arabidopsis thaliana] >sp BAA89784 BAA89784 IRE homolog 1 (fragment). Length = 1023	zal protein [Homo sapiens] L122098) hypothetical  -pir[T34532[T34532 KFZp434B1517.1 - human 266[CAB59266 Hypothetical ent). Length				similar to [SwissProt Accession Number P23842]; start codon is not identified yet [Escherichia coli] >sp P76954 P76954 YFEA PROTEIN (FRAGMENT). Length = 771			(AB005289) ABC transporter 7 protein [Homo sapiens] >sp[O75027]ABC7_HUMAN ATP-BINDING CASSETTE TRANSPORTER 7 PRECURSOR (ABC TRANSPORTER 7 PROTEIN). Length = 752
894341	894397	894631	894806	894811	894818	894820	894824	894827	894830
2044	2045	2046	2047	2048	2049	2050	2051	2052	2053

HCQDE22 Lambda ZAP	pSport1	pCMVSport 3.0	pSport1	HOEOQ19 Uni-ZAP XR	pSport1	HOUHL17 Uni-ZAP XR	pCMVSport 3.0	pSport1	pSport1	pSport1	Uni-ZAP XR
нсорб22	HWLVU33	HAJAY88	HCRPM46	ноеоо19	HKGBP52	HOUHL17	HDPPB40	HWL0I29	HCRMJ47	HLDXE66	HAIBM54
49				58	81				91	87	
48				39	9/				91	87	
841	388	3134	230	1049	1662	632	2310	552	1266	373	446
314	2	2856	3	3	1468	321	1873	337	112	2	117
6331	6332	6333	6334	6335	6336	6337	6338	6339	6340	6341	6342
gb AAC25416.  1				emb CAA7459 1.1	dbj BAA91131 				dbj BAA85892 .1	gb AAA60282. 1	
(AF072816) ABC-type transporter MRP3 [Rattus norvegicus] >sp 088563 MRP3_RAT CANALICULAR MULTISPECIFIC ORGANIC ANION TRANSPORTER 2 (MULTIDRUG RESISTANCE-ASSOCIATED PROTEIN 3) (MRP-LIKE PROTEIN-2) (MLP-2). Length = 1522				MAP3K delta-1 protein kinase [Arabidopsis thaliana] emb CAA7459 > sp O23719 O23719 MAP3K DELTA-1 PROTEIN 1.1  KINASE (FRAGMENT). Length = 406	(AK000385) unnamed protein product [Homo sapiens] Length = 152				(AB027466) spondin 2 [Homo sapiens]   dbj BAA85892   >sp BAA85892 BAA85892 Spondin 2. Length = 331   .1	ribosomal protein L7a large subunit [Homo sapiens] >emb CAA36383.1  L7a protein [Homo sapiens] >emb CAA29889.1  PLA-X polypeptide [Homo sapiens] >emb CAA43925.1  ribosomal protein L7a [Homo sapiens] >emb CAA33117.1  ribosomal protein L7a (AA 1-266) [Rattu	
894831	894832	894842	894878	895122	895303	895372	895675	895781	895927	800968	897234
2054	2055	2056	2057	2058	2059	2060	2061	2062	2063	2064	2065

2066	897524	unnamed protein product [Homo sapiens] >emb CAA93157.1  translocon-associated protein delta subunit precursor [Homo sapiens] >emb CAA92215.1  translocon-associated protein delta subunit precursor [Homo sapiens] >gb AAC51745.1  translocon-associated protein	emb CAA6221 1.1	6343	8	089	100	100	HSXAX45	HSXAX45 Uni-ZAP XR
2067	897898	ris] ) KD	emb CAB4586	6344	8	707	30	47	HE8PB56	Uni-ZAP XR
2068	898087	(AF072128) claudin-2 [Mus musculus] >sp[O88552 CLD2_MOUSE CLAUDIN-2. Length = 230	gb AAC27079.	6345	-	264	87	16	HTPGE66	HTPGE66 Uni-ZAP XR
5069	898136			6346	190	504			HWLIL19	pSport1
2070	898157	novel ORF [Homo sapiens] >sp O00251 O00251 HYPOTHETICAL PROTEIN (FRAGMENT). Length = 68	gb AAB72234.	6347	889	1002	8	88	HPJEE80	Uni-ZAP XR
2071	898192			6348	111	353			HWLQX67	pSport1
2072	898355	beta-galactosidase [Expression vector pBSII- LUCINT] Length = 69	gb AAB53629.	6349	1810	2109	94	86	HCRNK75	pSport1
2073	898418	(AC003965) SP001LA [Homo sapiens] Length = 271	gb AAB93671.  1	6350	2	694	82	84	HOGDR01	pCMVSport 2.0
2074	898427	reticulocalbin [Homo sapiens] >emb CAB53067.1  (AL078612) dJ65P5.1 (reticulocalbin 1, EF-hand calcium binding domain) [Homo sapiens] >pir JC4173 JC4173 reticulocalbin precursor - human >sp Q15293 RCN1_HUMAN RETICULOCALBIN 1 PRECURSOR. >sp CAB53067 CAB5306	dbj BAA07670 .1	6351	298	1017	64	78	HHATR06	pCMVSport 3.0
2075	898541	(AF169677) leucine-rich repeat transmembrane protein FLRT3 [Homo sapiens] >sp AAF28461 AAF28461 Leucine-rich repeat transmembrane protein FLRT3. Length = 649	gb AAF28461. 1 AF1696	6352	424	1938	06	06	нгормол	HLQDM07 Lambda ZAP

pCMVSport 3.0	pSport1	pBluescript SK-	Uni-ZAP XR	pSport1
HDPBW68	HISCJ15	HCYBH77	HPJAS61	HCRMK25
91	98	100	92	94
91	98	100	92	94
2761	2187	1656	1445	2630
605	_	-	m	1308
6353	6354	6355	6356	6357
gb AAD29125. 1 AF1264	gb AAD22032.	gb AAF35260. 1 AF0818	gb AAD45240. 1	sp[P02452 CA
(AF126484) CARD4 [Homo sapiens] >gb AAD28350.1 AF113925_1 (AF113925) Nod1 [Homo sapiens] >gb AAD43922.1  (AF149774) NOD1 protein [Homo sapiens] >sp Q9Y239 Q9Y239 NOD1 PROTEIN. Length = 953	(AF117754) thyroid hormone receptor-associated protein complex component TRAP240 [Homo sapiens] >sp AAD22032 AAD22032 Thyroid hormone receptor-associated protein complex component TRAP240. >dbj BAA25519.1  (AB011165) KIAA0593 protein [Homo sapiens] {SUB 1	(AF081886) ERO1-like protein [Homo sapiens] >gb AAF06104.1 AF123887_1 (AF123887) ERO1L [Homo sapiens] {SUB 84-468} Length = 468	(AF117892) aspartic-like protease [Homo sapiens] gb AAD45240. >gb AAD45963.1 AF050171_1 (AF050171) aspartyl 1  protease [Homo sapiens] >gb AAF17078.1  (AF200342) aspartyl protease 1 [Homo sapiens] >gb AAF26368.1 AF204944_1 (AF204944) transmembrane aspartic proteinase Asp	COLLAGEN ALPHA 1(1) CHAIN PRECURSOR. >emb CAA67261.1  collagen type I alpha 1 [Homo sapiens] {SUB 1-1069} >emb CAA29605.1  C-terminal propeptide domain [Homo sapiens] {SUB 1229-1454} Length = 1464
898651	898814	898946	899130	899224
2076	2077	2078	2079	2080

l .	899632	unnamed protein product [unidentified]  >pir A26359 A26359 decay-accelerating factor splice [0.1] form 1 precursor - human >dbj BAA22900.1  (AB003312) decay accelerating factor [Homo sapiens] {SUB 291-345} Length = 440	emb CAA0384	6358	ε	1376	87	87	HNTRV11	pSport1
	899644	lac repressor [Cloning vector pCMVLacl] >gb[AAC73448.1] (AE000141) transcriptional repressor of the lac operon [Escherichia coli] {SUB 1-360} >gb[AAB61949.1] lac repressor fragment [unidentified cloning vector] {SUB 331-360} >gb[AAA72793.1] lacl [unidenti	gb AAB17268.	6359	г.	401	96	97	HWLOU33	pSport1
1	899661	(AL050297) hypothetical protein [Homo sapiens] >emb CAB43396.1  (AL050297) hypothetical protein [Homo sapiens] >pir T08701 T08701 hypothetical protein DKFZp564N123.1 - human (fragment) >sp Q9Y3T6 Q9Y3T6 HYPOTHETICAL 50.0 KD PROTEIN (FRAGMENT). Length = 45	emb CAB4339 6.1	6360	21	1016	06	91	HAPNOS0	HAPNOS0 Uni-ZAP XR
1	922668	formate dehydrogenase [Escherichia coli] Length = 715	gb AAA23754.	6361	619	1392	66	66	HBSAK60	HBSAK60 Uni-ZAP XR
l .	998668	artifact-warning sequence (translated ALU class F) - human Length = $673$	pir F40201 F40 201	6362	88	201	59	70	HDPOD73	pCMVSport 3.0
	899885	SDF2 [Homo sapiens] >pirJC5106 JC5106 stromal cell-derived factor 2 precursor - human >sp Q99470 Q99470 SDF2. Length = 211	dbj BAA09312 .1	6363	233	928	91	92	НWННQ57	pCMVSport 3.0
	899913	(AB012223) ORF2 [Canis familiaris] >sp O62658 O62658 LINE-1 ELEMENT ORF2. Length = 1275	dbj BAA25253 .1	6364	238	92	36	59	HNFHY51	Uni-ZAP XR
1	900015			6365	1	222			HTOHV42	HTOHV42 Uni-ZAP XR
1	900162	(AF157106) soluble secreted endopeptidase delta [Mus musculus] >sp AAF13153 AAF13153 Soluble secreted endopeptidase delta. Length = 742	gb AAF13153. 1 AF1571	6366	367	753	82	94	HWLXO02	pSportl

				+	+	
pSport1	pSport1	pSport1	pSport1	pCMVSport 1	pCMVSport 3.0	pBluescript SK-
HWLKM77	HWMCJ06	HCRPZ48	HCRMU04	ннвеа82	HWHGX93 pCMVSport 3.0	HTNAI80
93	76	57	84	95	26	66
93	76	47	84	94	<i>L</i> 6	66
1409	2143	902	1492	3081	1132	1328
m	29	21	611	250	2	£
6367	6368	6369	6370	6371	6372	6373
gb AAD15624.	emb CAB6374 6.1	gb AAF03695. 1 AF1755	dbj BAA13385 .1	gb AAF07395. 1 AF1060	pir JE0174 JE0 174	gb AAD25870. 1 AF0207
(AC006950) IgG Fc binding protein [AA 4671-5405] gb[AAD15624. [Homo sapiens] >sp[O95784 O95784 IGG FC 1  BINDING PROTEIN (FRAGMENT). Length = 735	(AL133620) hypothetical protein [Homo sapiens] >emb CAB63746.1  (AL133620) hypothetical protein [Homo sapiens] >sp CAB63746 CAB63746 Hypothetical 115.3 kd protein. Length = 1031	(AF175522) transmembrane tryptase [Homo sapiens] gb AAF03695. >gb AAF03697.1 AF175759_1 (AF175759)	Similar to S.cerevisiae EMP70 protein precursor (S25110) [Homo sapiens] >sp Q92544 Q92544 MYELOBLAST KIAA0255. Length = 625	(AF106037) adipocyte-derived leucine aminopeptidase [Homo sapiens] >sp AAF07395 AAF07395 Adipocyte-derived leucine aminopeptidase. Length = 941	frizzled protein-2 - human Length = 295	(AF020797) AP-mu chain family member mu1B [Homo sapiens] >splQ9Y6Q5[Q9Y6Q5 AP-MU CHAIN FAMILY MEMBER MU1B. Length = 423
900249	900555	969006	900777	900784	900838	900919
2090	2091	2092	2093	2094	2095	2096

1	996006		emb CAB5596	1	1784	2824	64	<del></del>	HCRPO45	pSport1
900991		(AF077030) hypothetical 43.2 kDa protein [Homo gb AAD2 sapiens] >gb AAD34049.1 AF151812_1 (AF151812) 1 AF0770 CGI-54 protein [Homo sapiens] >sp Q9Y282 Q9Y282 CGI-54 PROTEIN. Length = 383	gb AAD27763. 1 AF0770	6375	-	1230	96	96	HWLWF60	pSport1
006	900993	112 [Homo 78 Transmembrane 5	gb AAD55678. 1 AF1477	6376	3	1619	84	98	HCNCY58	HCNCY58 Lambda ZAP
901111	111		gb AAD17522. 1	6377	3	1097	84	85	HCNDA61	HCNDA61 Lambda ZAP
901125	125	(AF127036) calcium-activated chloride channel protein 1 [Homo sapiens] >sp AAD25487/AAD25487 Calcium-activated chloride channel protein 1. Length = 914	gb AAD25487. 1 AF1270	6378	23	2629	96	96	HCNUB65	pBluescript
901128	128	(AB002362) KIAA0364 [Homo sapiens] >gb AAC52057.1  (AF034198) IGSF1 [Homo sapiens] >pir T09402 T09402 immunoglobulin-like protein IGSF1 - human >sp O15070 O15070 KIAA0364. Length = 1327	dbj BAA20819	6379	2	826	42	57	HWLRB02	pSport1
901202	202	(AJ245620) CTL1 protein [Homo sapiens] Length = 654	emb CAB7554 1.1	6380	98	2209	93	93	HSDKL35	Uni-ZAP XR
901253	253	GallNAc alpha-2,6-sialyltransferase I [Mus musculus] emb CAA7213 > sp CAA72137 CAA72137 GallNAc alpha-2,6-sialyltransferase I. Length = 526	emb CAA7213 7.1	6381	31	1905	9	73	HJPCX37	Uni-ZAP XR
901	901276	(AB023172) KIAA0955 protein [Homo sapiens] >sp Q9Y2G2 Q9Y2G2 KIAA0955 PROTEIN. Length = 431	dbj BAA76799 .1	6382	259	1155	86	66	HPBEM10	pBluescript SK-

pCMVSport 3.0	pBluescript SK-	Lambda ZAP II	Uni-ZAP XR	Uni-ZAP XR	Lambda ZAP II
HWBDL33	H2LBA47	нсол172	нетнс61	HTXLJ25	HCNAI22
49	09	100	46	71	80
35	36	100	29	52	62
1288	699	209	819	724	671
470	172	2	<i>L</i> 9	2	8
6383	6384	6385	6386	6387	6388
gb AAF17243. 1 AF2019	4bj BAA03111	gb AAC77358. 1	gb AAF02448. 1 AF1255	emb CAA9299	emb CAA6064 5.1
(AF201951) high affinity immunoglobulin epsilon receptor beta subunit [Homo sapiens] Length = 240	regenerating protein I [Mus musculus]  >pir A47148 A47148 reg I, regenerating islet cells - mouse >sp P43137 LIT1_MOUSE  LITHOSTATHINE I PRECURSOR (PANCREATIC STONE PROTEIN I) (PSP) (PANCREATIC THREAD PROTEIN I) (PTP) (ISLET OF LANGERHANS REGENERATING PROTE	(AF007791) secreted cement gland protein XAG-2 homolog [Homo sapiens] >gb[AAC82614.1] (AF038451) secreted cement gland protein XAG-2 homolog [Homo sapiens] >gb[AAF22484.1 AF088867_1 (AF088867) putative secreted protein XAG [Homo sapiens] >pir[JE0350]JE035	(AF125543) major histocompatibility complex class I protein [Monodelphis domestica] >sp AAF02448 AAF02448 Major histocompatibility complex class I protein. Length = 347	predicted using Genefinder; Similarity to Mouse FK506-binding protein (SW:FKB3_MOUSE) [Caenorhabditis elegans] >pir[T21882[T21882] hypothetical protein F36H1.1 - Caenorhabditis elegans >sp Q20107 Q20107 F36H1.1 PROTEIN. Length = 139	2.19 [Homo sapiens] >emb CAA39090.1  2-19 protein [Homo sapiens] >gb AAA92652.1  2_19 [Homo sapiens] >pir I37095 I37095 gene 2.19 protein - human >sp P98173 219_HUMAN 2-19 PROTEIN PRECURSOR. Length = 230
901333	901375	901415	901421	901472	901473
2106	2107	2108	2109	2110	2111

2112	901494	(AB006781) galectin-4 [Homo sapiens] >gb AAB86590.1  galectin-4 [Homo sapiens] >gb AAC51763.1  (AF014838) galectin-4 [Homo sapiens] >sp F56470 LEG4_HUMAN GALECTIN-4 (LACTOSE-BINDING LECTIN 4) (L-36 LACTOSE BINDING PROTEIN) (L36LBP). >sp AAB86590 AAB86590	dbj BAA22165 .1	6389	2	805	92	92	HSIAL77	HSIAL77 Uni-ZAP XR
2113	901515	pre-pump-1 proteinase (AA -17 to 250) [Homo sapiens] >emb CAA77942.1  PUMP [Homo sapiens] >pir B28816 KCHUM matrilysin (EC 3.4.24.23) precursor - human >sp P09237 COG7_HUMAN MATRILYSIN PRECURSOR (EC 3.4.24.23) (PUMP-1 PROTEASE) (UTERINE METALLOPROTEINASE)	emb CAA3067 8.1	6390	<u>س</u>	839	100	100	HRACJ32	pCMVSport 3.0
2114	901567	(AK001466) unnamed protein product [Homo sapiens] Length = 202	dbj BAA91708 .1	6391	2	736	92	92	HMGBJ25	HMGBJ25 Uni-ZAP XR
2115	901578	ż.	emb CAB0214 3.1	6392	2	1417	08	08	HDTE010	pCMVSport 2.0
2116	901621	(AL031685) d1963K23.2 (novel protein) [Homo sapiens] >sp Q9Y508 Q9Y508 DJ963K23.2 (NOVEL PROTEIN) (FRAGMENT). Length = 228	emb CAB4602 8.1	6393	2	694	94	94	HSSGC06	HSSGC06 Uni-ZAP XR
2117	901875	GalNAc alpha-2,6-sialyltransferase I [Mus musculus] emb CAA7213 >sp CAA72137 CAA72137 GalNAc alpha-2,6- 7.1  sialyltransferase I. Length = 526	emb CAA7213 7.1	6394	32	1906	<i>L</i> 9	75	HSICN14	Uni-ZAP XR
2118	HWLM073R	HWLMO73R !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! sp P39194 AL Length = 593	sp P39194 AL U7_HUMAN	6395	2	496	81	98	HWLMO73	pSport1
2119	HCRMU56R	HCRMU56R   !!!! ALU SUBFAMILY SX WARNING ENTRY !!!!   sp P39195 AL   Length = 591	sp P39195 AL U8_HUMAN	9689	25	129	75	77	HCRMU56	pSport1

pSport1	pBluescript	pSport1	HCQCG26 Lambda ZAP	pSport1	pSport1	Uni-ZAP XR	Uni-ZAP XR
HWLRH49	HKCSA70	HWLOB10	нсос626	HCRNR57	HWLUZ40	H6EBJ04	HOENF69
75	100	95	86	95	62		46
75	100	93	86	95	79		43
115	159	163	219	303	313	129	264
2	_	2		1	2	332	-
6397	6398	6399	6400	6401	6402	6403	6404
dbj BAA03604	dbj BAA19211	dbj BAA22388 .1	dbj BAA20764 .1	dbj BAA20765 .1	gb AAD43978. 1 AF1529		dbj BAA25534 .1
HWLRH49R "TFIIA-42" [Homo sapiens] >emb CAA53151.1  TFIIA [Homo sapiens] >emb CAA54442.1  TFIIA/alpha, p55 [Homo sapiens] >gb AAF26776.1 AC010582_2 (AC010582) TFIIA- 42 [Homo sapiens] >gb AAF26776.1 AC010582_2 (AC010582) TFIIA-42 [Homo sapiens] >pir A49077 A49077	(AB000911) ribosomal protein [Sus scrofa] >emb CAB56794.1  ribosomal protein S18 [Homo sapiens] >emb CAA20231.1  (AL031228) dJ1033B10.4 (40S ribosomal protein S18 (RPS18, KE-3)) [Homo sapiens] >gb AAA16795.1  ribosomal protein [Mus musculus] >emb CAA40750	(AB001428) motor domain of KIF12 [Mus musculus] >sp 035061 035061 MOTOR DOMAIN OF KIF12 (FRAGMENT). Length = 169	(AB002303) KIAA0305 [Homo sapiens] >sp O15023 O15023 KIAA0305. Length = 1539	(AB002304) KIAA0306 [Homo sapiens] >sp BAA20765 BAA20765 KIAA0306 protein (fragment). Length = 1451	HWLUZ40R (AF152961) chromatin-specific transcription elongation factor FACT 140 kDa subunit [Homo sapiens] >sp Q9Y5B9 Q9Y5B9 CHROMATIN-SPECIFIC TRANSCRIPTION ELONGATION FACTOR FACT 140 KDA SUBUNIT. >gb AAF28231.1  (AF164924) SPT16/CDC68-like protein [Homo sapiens]		(AB011180) KIAA0608 protein [Homo sapiens] >sp O60347 O60347 KIAA0608 PROTEIN (FRAGMENT). Length = 775
HWLRH49R	HKCSA70R	HWLOB10R	HCQCG26R	HCRNR57R	HWLUZ40R	H6EBJ04R	HOENF69R
2120	2121	2122	2123	2124	2125	2126	2127

Lambda ZAP II	pSport1	pSport1	pBluescript	Lambda ZAP II	pSport1	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-
нсорс81	HWLQY33	HCRNF08	HKCSZ69	HCQAG23	<b>НС</b> КQD03	H2LAF75	H2LAJ65	H2LAT73	H2LAX35
08	95	82	86	94	75	96		96	93
92	95	79	86	92	99	96		96	93
393	382	145	313	295	573	180	159	202	150
169	197	2	2	149	1	19	1	2	-
6405	6406	6407	6408	6409	6410	6411	6412	6413	6414
dbj BAA31522 .1	dbj BAA31594 .1	pir T00358 T0 0358	gb AAA36597.	dbj BAA34512 .1	gb AAA69736. 1	sp Q9Y6Y5 Q9 Y6Y5		sp Q9Y6Y5 Q9 Y6Y5	emb CAA0647 1.1
HCQDC81R (AB012725) zinc finger protein [Mus musculus] >sp O88291 O88291 ZINC FINGER PROTEIN. Length = 580	KIAA0619 protein [Homo sapiens] 075116 KIAA0619 PROTEIN. Length	HCRNF08R hypothetical protein KIAA0684 - human (fragment)   >sp 075169 075169 KIAA0684 PROTEIN (FRAGMENT). Length = 903	scar protein [Homo sapiens] Length = 244	(AB018335) KIAA0792 protein [Homo sapiens] >sp O94886 O94886 KIAA0792 PROTEIN. Length = 807	HCRQD03R immunoglobulin heavy chain [Homo sapiens] Length gb AAA69736.	IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414		IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	(AJ005324) glutamate permease [synthetic construct] emb CAA0647 >emb CAA06474.1  (AJ005327) glutamate permease [1.1] [synthetic construct] >emb CAA06477.1  (AJ005330) glutamate permease [synthetic construct] >eb AAA24514.1  gltS [Escherichia coli] {SUB 437-459} Length = 45
HCQDC81R	HWLQY33R	HCRNF08R	HKCSZ69R	HCQAG23R	HCRQD03R	H2LAF75R	H2LAJ65R	H2LAT73R	H2LAX35R
2128	2129	2130	2131	2132	2133	2134	2135	2136	2137

pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	HTXPR08 Uni-ZAP XR
H2LAX79	HCYBK85	нсувк96	HCYBL18	HCYBM62	HCYBO61	HTXPR08
96		87	98	08	100	100
96		87	08	75	100	100
158	127	154	120	123	112	88
m	11	2	_	-	2	2
6415	6416	6417	6418	6419	6420	6421
sp Q9Y6Y5 Q9 Y6Y5		\$p Q9Y6Y5 Q9 Y6Y5	\$p Q9Y6Y5 Q9 Y6Y5	sp Q9Y6Y5 Q9 Y6Y5	\$p Q9Y6Y5 Q9 Y6Y5	dbj BAA74886 .1
H2LAX79R IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414		IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	IDN4-GGTR14 PROTEIN >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	₹	HCYBO61R IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	HTXPR08R (AB020670) KIAA0863 protein [Homo sapiens] >sp O94943 O94943 KIAA0863 PROTEIN. Length = 1131
H2LAX79R	HCYBK85R	HCYBK96R	HCYBL18R	HCYBM62R	HCYB061R	HTXPR08R
2138	2139	2140	2141	2142	2143	2144

pSportl	251 pSport1	HKLAB44 Lambda ZAP	.06 pBluescript SK-	.41 pSport1	HCNAH60 Lambda ZAP	'58 Lambda ZAP II
HUUAQ45	HWLWQ51	HKLAB	H2CBA06	HWLRL41	HCNAH	HCNDF58
78	9/	84	100	100	81	98
78	73	82	100	94	08	83
322	517	389	354	154	325	328
89	2	<del>د</del>	-	2	2	62
6422	6423	6424	6425	6426	6427	6428
.1	dbj BAA76818 .1	dbj BAA76837 .1	dbj BAA76849 .1	dbj BAA78778 .1	gb AAC08708. 1	gb AAC08708. 1
HUUAQ45R (AB021288) beta 2-microglobulin [Homo sapiens] >gb AAA87972.1  beta-2-microglobulin [Pan troglodytes] >gb AAA88008.1  beta-2-microglobulin [Gorilla gorilla] >gb AAD48083.1  (AF072097) beta- 2 microglobulin [Homo sapiens] >pir A90976 MGHUB2 beta-2-microglob	HWLWQ51R (AB023191) KIAA0974 protein [Homo sapiens] >sp Q9Y210 Q9Y210 KIAA0974 PROTEIN (FRAGMENT). Length = 565	in [Homo sapiens] 93 PROTEIN		nit e NIT E	HCNAH60R (AC002301) Homolog of rat Zymogen granule membrane protein [Homo sapiens] >sp O60844 O60844 HOMOLOG OF RAT ZYMOGEN GRANULE MEMBRANE PROTEIN. Length = 167	(AC002301) Homolog of rat Zymogen granule membrane protein [Homo sapiens] >sp O60844 O60844 HOMOLOG OF RAT ZYMOGEN GRANULE MEMBRANE PROTEIN. Length = 167
HUUAQ45R	HWLWQ51R	HKLAB44R	H2CBA06R	HWLRL41R	HCNAH60R	HCNDF58R
2145	2146	2147	2148	2149	2150	2151

2152	HCRMZ60R	HCRMZ60R (AC002483) putative product from mRNA sequence gb AAC35295. CG003 from BRCA2 region; match to U50534 (NID:g1685103) [Homo sapiens] >pir[T00415 T00415 hypothetical protein H_248015.1 - human (fragment) >sp 014572 014572 WUGSC:H_248015.1 PROTEIN (FRAGMENT). Length = 184	gb AAC35295.	6429	2	544	29	67	HCRMZ60	pSport1
2153	HCRNL13R	(AC002563) putative RHO/RAC effector protein; 95% similarity to P49205 (PID:g1345860) [Homo sapiens] >sp[014578 CTRO_HUMAN CITRON PROTEIN (FRAGMENT). >dbj BAA76793.1  (AB023166) KIAA0949 protein [Homo sapiens] {SUB 347-1286} Length = 1286	gb AAB71327.	6430	۳	, ,	86	100	HCRNL13	pSport1
2154	HCRMX17R	HCRMX17R (AC003682) R27945_2 [Homo sapiens] >sp O43338 O43338 R27945_2. Length = 475	gb AAC24607.	6431	19	204	51	9	HCRMX17	pSport1
2155	HWMBJ68R	6.1 (novel protein) [Homo 791 CAB65791 DJ842G6.1 nent). Length = 197	emb CAB6579 1.1	6432	3	218	001	100	HWMBJ68	pSport1
2156	HWLOV91R	HWLOV91R (AC004475) F23858_1 [Homo sapiens] >pir[T02299 T02299 hypothetical protein F23858_1 - human (fragment) >splO60378 O60378 F23858_1   (FRAGMENT). >emb CAB70678.1  (AL137286) hypothetical protein [Homo sapiens] {SUB 217-608} Length = 608	gb AAC08052. 1	6433	3	455	99	99	HWLOV91	pSport1
2157	HCRPW27R	HCRPW27R (AC004528) R32184_3 [Homo sapiens] >sp 060392 060392 R32184_3. Length = 529	gb AAC12681.	6434	3	314	96	96	HCRPW27	pSport1
2158	HELGR96R	APP-binding protein 1 [Rattus norvegicus] >sp Q9Z1A5 Q9Z1A5 APP-BINDING PROTEIN 1. Length = 534	gb AAD09247. 1	6435	1	249	75	62	HELGR96	Uni-ZAP XR
2159	HCRPB14R			6436	2	79			HCRPB14	pSport1
2160	HCRQM72R			6437	273	1			HCRQM72	pSport1
2161	HWLNK47R			6438	343	2			HWLNK47	pSport1

pSport1	pSport1	pBluescript SK-	pSport1	nbda ZAP II	pSport1	pSport1
HWLOI40 F	100 HWLMH52 F	H2CBU03 pB	HWLUL19 F	HCQDR91 Lambda ZAP	100 HWMBN34 F	HWMBS08
93	100	96	62 1	06	100 F	74 I
98	100	96	59	06	100	65
212	245	353	211	146	388	426
403	r.	3	2	385	2	22
6439	6440	6441	6442	6443	6444	6445
emb CAB6919 5.1	emb CAB5140 5.1	dbj BAA86532 .1	gb AAC23790.	emb CAB5600 5.1	gb AAD15550. 1	gb AAD23607. 1 AC0072
HWLOI40R unnamed protein product [unidentified] Length = 180 emb CAB6919   5.1	HWLMH52R (AL096881) hypothetical protein [Homo sapiens] >gb AAF19256.1 AC004832_1 (AC004832) similar to 45 kDa secretory protein [Rattus norvegicus]; similar to CAA10644.1 (PID:g4164418) [Homo sapiens] >sp O76054 O76054 HYPOTHETICAL 46.1 KD PROTEIN. >sp AAF19256 A	(AB033044) KIAA1218 protein [Homo sapiens] >sp BAA86532 BAA86532 KIAA1218 protein (fragment). Length = 864	HWLUL19R (AC005154) similar to protein U28928 (PID:g861306) [Homo sapiens] >sp[075223 075223 WUGSC:H_D10777023.1 PROTEIN. Length = 188	HCQDR91R (AL117583) hypothetical protein [Homo sapiens] >emb CAB56005.1  (AL117583) hypothetical protein [Homo sapiens] >pir[T17315 T17315] hypothetical protein DKFZp434K058.1 - human >sp CAB56005 CAB56005 Hypothetical 12.7 kd protein. Length = 117	HWMBN34R (AC006153) similar to Aquifex aeolicus GTP- binding protein; similar to AE000771 (PID:g2984292) [Homo sapiens] >splQ9Y6T6[Q9Y6T6 WUGSC:H_NH0120J02.1 PROTEIN (FRAGMENT). Length = 206	HWMBS08R (AC007228) BC37295_1 [Homo sapiens] >sp[Q9Y2N9 Q9Y2N9 BC37295_1. Length = 599
HWLOI40R	HWLMH52R	H2CBU03R	HWLUL19R	нсоркогк	HWMBN34R	HWMBS08R
2162	2163	2164	2165	2166	2167	2168

pBluescript	pSport1	pCMVSport 2.0	pBluescript	pSport1	Lambda ZAP II	pBluescript SK-
HKCSC14	HCRNF81	НОНСІЗ1	HSKKC10	HWLWU01	нсорѕ79	H2CBC52
70	06	62	77	95	92	
89	88	57	72	95	85	
41	591	327	213	246	571	370
427	-	521	34	1	08	50
6446	6447	6448	6449	6450	6451	6452
gb AAC74378.	gb AAC39540. 1	gb AAB81938. 1	gb AAB64189. 1	gb AAB87763. 1	gb AAC19158. 1	
(AE000227) putative amino acid/amine transport protein [Escherichia coli] >pir C64878 C64878 probable amino acid permease ycjJ - Escherichia coli >sp P76037 YCJJ ECOLI HYPOTHETICAL 50.9 KD TRANSPORT PROTEIN IN SAPA-ALDH INTERGENIC REGION. {SUB 19-479} >db	(AF000364) heterogeneous nuclear ribonucleoprotein R [Homo sapiens] >pir[T02673[T02673 heterogeneous nuclear ribonucleoprotein R - human >sp O43390 O43390 HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN R. Length = 633	(AF000381) non-functional folate binding protein [Homo sapiens] >sp O14597 O14597 NON-FUNCTIONAL FOLATE BINDING PROTEIN. Length = 254	(AF006084) p41-Arc [Homo sapiens] >gb AAF03508.1 AC004922_5 (AC004922) P41- ARC [Homo sapiens] >sp O15143 AR41_HUMAN ARP2/3 COMPLEX 41 KD SUBUNIT (P41-ARC). >sp AAF03508 AAF03508 P41-ARC. Length = 372	<u>S</u>	(AF007157) [Homo sapiens] >sp 043414 043414 HYPOTHETICAL 43.1 KD PROTEIN (FRAGMENT). >gb AAC04618.1  (AC004254) similar to C. elegans hypothetical protein; similar to AF038615 (PID:g2736329) [Homo sapiens] {SUB 224-312} >emb CAB55430.1  (AL035417)	
HKCSC14R	HCRNF81R	HOHCI31R	HSKKC10R	HWLWU01R	нсдрѕ79к	H2CBC52R
2169	2170	2171	2172	2173	2174	2175

176	HWLWU22R	2176 HWLWU22R (AF012023) integrin cytoplasmic domain associated [gb AAB88671. protein; Icap-1a [Homo sapiens]   1    >sp O14713 O14713 INTEGRIN CYTOPLASMIC   DOMAIN ASSOCIATED PROTEIN. Length = 200	gb AAB88671. 1	6453		342	88	88	HWLWU22	pSport1
2177	HWLMC24R	HWLMC24R (AF051100) immunoglobulin G Fd fragment [Homo gb AAD15787. sapiens] Length = 223	gb AAD15787.	6454	82	366	77	77	HWLMC24	pSport1
2178	HWLUR40R	e-associated kinase [Homo 14731 MEMBRANE-SE. Length = 499	gb AAB71843. 1	6455	_	324	88	06	HWLUR40	pSport1
2179	нсосн96к			6456	94	156			96НОООН	HCQCH96 Lambda ZAP
2180	HHAOD46R	HHAOD46R (AF014955) TFAR19 [Homo sapiens] >pir JG0192 JG0192 TFAR19 protein - human >sp O14737 TF19 HUMAN TFAR19 PROTEIN (TF-1 CELL APOPTOSIS RELATED GENE-19 PROTEIN). Length = 125	gb AAD11579.	6457	49	432	85	88	ННАОД46	pCMVSport 3.0
2181	HCYBA83R	(AF017061) vasopressin-activated calcium mobilizing putative receptor protein [Homo sapiens] Length = 781	gb AAB70253.	6458	_	489	86	66	HCYBA83	pBluescript SK-
2182	HCROZ77R	HCROZ77R (AF018261) EH domain binding protein Epsin [Rattus norvegicus] >sp[O88339]O88339 EH DOMAIN BINDING PROTEIN EPSIN. Length = 575	gb AAC33823. 1	6459	1	525	<i>L</i> 9	69	HCROZ77	pSport1
2183	HWLND06R	HWLND06R (AF022108) putative replication initiator origin recognition complex subunit Orc4Lp [Homo sapiens] >gb AAC80282.1  (AF047598) origin recognition complex subunit 4; Orc4p [Homo sapiens] >gb AAD22110.1  (AF132596) origin recognition complex subunit 4 [Homo	gb AAC01957.	6460	-	213	73	79	HWLND06	pSport1
2184	HCQCP20R	(AB038463) GC36 [Homo sapiens] Length = 664	dbj BAA92137 .1	6461	2	229	86	86	нсось20	HCQCP20 Lambda ZAP

HCRQF71 pSport1	HWLNF84 pSport1	HCRMF28 pSport1	HCROQ32 pSport1	100 HWLOW79 pSport1	HCRQL67 pSport1	HCRQI10 pSport1	HWLRE34 pSport1
100	52	08	96		17	66	100
100	33	74	06	100	89	66	92
383	457	184	297	133	551	299	242
<i>د</i>	2	∞	-	2	21	٤	153
6462	6463	6464	6465	6466	6467	6468	. 6469
gb AAB80762.	gb AAB70984. 1	gb AAB82609. 1	gb AAC32547. 1	gb AAC18823. 1	gb AAB86408. 1	emb CAB7073 6.1	gb AAC26103. 1
HCRQF71R (AF022799) digestive tract-specific calpain; calcium- gb AAB80762. dependent cysteine proteinase [Homo sapiens]	HWLNF84R (AF025459) H14A12.3 gene product [Caenorhabditis gb AAB70984. elegans] >sp O17213 O17213 H14A12.3 PROTEIN. 1  Length = 284	(AF026977) microsomal glutathione S-transferase 3 [Homo sapiens] >sp O14880 O14880 MICROSOMAL GLUTATHIONE S-TRANSFERASE 3. Length = 152	HCROQ32R (AF029789) GTPase-activating protein [Homo sapiens] > sp O60484 O60484 GTPASE-ACTIVATING PROTEIN. Length = 1041	HWLOW79R (AF030339) VESPR [Homo sapiens] >pir T09074 T09074 semaphorin receptor VESPR - human >sp O60486 O60486 VESPR. Length = 1568	HCRQL67R (AF030430) semaphorin VIa [Mus musculus] >sp O35464 O35464 SEMAPHORIN VIA PRECURSOR. Length = 888	(AL137438) hypothetical protein [Homo sapiens] >emb CAB70736.1  (AL137438) hypothetical protein [Homo sapiens] >sp CAB70736 CAB70736 Hypothetical 69.3 kd protein (fragment). Length = 596	HWLRE34R (AF034802) liprin-beta1 [Homo sapiens] >sp 075336 075336 LIPRIN-BETA1. Length =
HCRQF71R	HWLNF84R	HCRMF28R	HCROQ32R	HWLOW79R	HCRQL67R	HCRQ110R	HWLRE34R
2185	2186	2187	2188	2189	2190	2191	2192

pSport1	pSport1	pSport1	HCQAI71 Lambda ZAP	HCQCQ75 Lambda ZAP II
HULCD94	HHMMF84	HCRP008	HCQAI71	нсосо75
98	100	100	<i>L</i> 9	93
82	100	94	41	88
466	113	267	289	298
119	m	yeard	2	2
6470	6471	6472	6473	6474
gb AAC39252.	gb AAC68838.	gb AAC39606.	gb AAB91418. 1	emb CAA4294
HULCD94R (AF035178) elongation factor 1 A2 [Oryctolagus cuniculus] >emb CAA50280.1  elongation factor 1 alpha-2 [Homo sapiens] >pir S35033 EFHUA2 translation elongation factor eEF-1 alpha-2 chain - human >sp Q05639 EF12_HUMAN ELONGATION FACTOR 1-ALPHA 2 (EF-1-ALPH	(AF035840) NADH:ubiquinone oxidoreductase B17 subunit [Homo sapiens] >gb AAD32451.1 AF067167_1 (AF067167) NADH-ubiquinone oxidoreductase B17 subunit homolog [Homo sapiens] >sp AAD32451 AAD32451 NADH-ubiquinone oxidoreductase B17 subunit homolog. >sp O9513	HCRP008R (AF035940) similar to mago nashi [Homo sapiens]  >gb AAD32457.1 AF067173_1 (AF067173) Mago homolog [Homo sapiens] >gb AAB66722.1  (AF007862) mm-Mago [Mus musculus]  >gb AAC40044.1  (AF035939) similar to mago nashi [Mus musculus] >sp P50606 MGN_HUMAN MAGO NA	HCQAI71R (AF037168) DnaJ homologue [Arabidopsis thaliana] >sp O49070 O49070 DNAJ HOMOLOGUE. Length = 284	HCQCQ75R polyubiquitin [Cricetulus griseus]  >pir S21083 S21083 polyubiquitin 5 - Chinese hamster >dbj BAA03983.1  polyubiquitin [Rattus norvegicus] (SUB 77-381) >gb AAA49129.1  ubiquitin polyprotein (heat shock related) [Gallus gallus] {SUB 225-381} Length = 381
HULCD94R	HHMMF84R	HCRPO08R	HCQAI71R	HCQCQ75R
2193	2194	2195	2196	2197

4 pSport1	pBluescript	pSport1	pBluescript SK-	3 pSport1	3 Lambda ZAP II	HCQDD24   Lambda ZAP	5 pCMVSport 2.0	7 pSport1	3 pSport1
HWLMQ74	HFVKA48	HLXNF14	H2LAB80	HOCTC23	нсорозз	нсорр2	HKAFL06	HCRMO57	HCRO083
76	95	100	68		8	53	83	63	97
73	95	100	68		83	20	78	59	6
281	247	144	476	208	188	376	465	330	250
06	2	-	3	2	ω	161	112	43	2
6475	6476	6477	6478	6479	6480	6481	6482	6483	6484
gb AAC03122.	gb AAD32452. 1 AF0671	gb AAC03787. 1	gb AAD04814. 1		gb AAC36629. 1	dbj BAA91179 .1	gb AAD17531. 1	emb CAB6943 4.1	gb AAD08657.
HWLMQ74R (AF047471) mitotic checkpoint protein kinase [Homo sapiens] >gb[AAC12729.1  (AF046078) protein kinase [Homo sapiens] >gb[AAD43675.1  (AF139363) BUB1 protein [Homo sapiens] >gb[AAB97855.2  (AF043294) putative mitotic checkpoint kinase [Homo sapiens] >sp[O4	(AF067168) NADH-ubiquinone oxidoreductase B22 subunit homolog [Homo sapiens] >sp AAD32452 AAD32452 NADH-ubiquinone oxidoreductase B22 subunit homolog. Length = 179	(AF047470) malate dehydrogenase precursor [Homo gb AAC03787. sapiens] >sp P40926 MDHM_HUMAN MALATE 1  DEHYDROGENASE, MITOCHONDRIAL PRECURSOR (EC 1.1.1.37). Length = 338	(AF047711) nGAP [Homo sapiens] >sp[095174 095174 NGAP. >emb CAB53260.1  (AL035702) dJ593C16.1 (ras GTPase activating protein) [Homo sapiens] {SUB 41-1139} Length = 1139		(AF052788) immunoglobulin light chain variable region [Homo sapiens] Length = 116	(AK000460) unnamed protein product [Homo sapiens] Length = 284	rotein [Homo sapiens] STAUFEN PROTEIN. F061938) staufen protein 82-577} Length = 577	unnamed protein product [unidentified] Length = 99	(AF071172) HERC2 [Homo sapiens] >sp 095714 095714 HERC2. Length = 4834
HWLMQ74R	HFVKA48R	HLXNF14R	H2LAB80R	HOCTC23R	нсороззк	HCQDD24R	HKAFL06R	HCRMO57R	HCROO83R
2198	2199	2200	2201	2202	2203	2204	2205	2206	2207

pSport1	pSport1	pSport1	pBluescript SK-	Uni-ZAP XR	pBluescript SK-	pSport1
	bSp	dSq	pBlu S	Uni-Z	pBlu S	i <sub>Sd</sub>
HCRMW16	HWL0035	HWĽVL <i>77</i>	HCYBH84	HBJMG15	н2Свн29	HWLMT35
	HW					
71		95	94	100	85	88
09		95	93	100	69	74
284	108	238	408	369	459	388
3	1	32	52	106	-	2
1	6486	6487	6488	6489	6490	6491
dbj BAA83042 .1		gb AAC63516. 1	gb AAD23915. 1	gb AAC83664. 1	gb AAC79841. 1	emb CAB3716
HCRMW16R (AB029013) KIAA1090 protein [Homo sapiens] >sp BAA83042 BAA83042 KIAA1090 protein. Length = 713		(AF073298) small EDRK-rich factor 2 [Homo sapiens] >gb AAC63515.1  (AF073297) small EDRK-rich factor 2 [Mus musculus] >sp O75918 O75918 4F5REL. >sp O88891 O88891 4F5REL. Length = 59	HCYBH84R (AF080000) RSK-like protein kinase RLPK [Homo sapiens] >gb AAC31171.1  (AF074393) nuclear mitogen- and stress-activated protein kinase-1 [Homo sapiens] >pir T13149 T13149 mitogen-and stress-activated protein kinase-1, nuclear - human >sp O75582 O75582 NUC	(AF080683) PITSLRE protein kinase alpha SV9 isoform [Homo sapiens] >sp 095265 095265 PITSLRE PROTEIN KINASE ALPHA SV9 ISOFORM. Length = 755	H2CBH29R (AF082556) TRF1-interacting ankyrin-related ADP-ribose polymerase [Homo sapiens] >sp 095271 095271 TRF1-INTERACTING ANKYRIN-RELATED ADP-RIBOSE POLYMERASE. >gb AAC79842.1 (AF082557) TRF1-interacting ankyrin-related ADP-ribose polymerase [Homo sapiens] {SU	HWLMT35R (AJ239373) immunoglobulin heavy chain variable region [Homo sapiens] >emb CAB37166.1  (AJ239373) immunoglobulin heavy chain variable region [Homo sapiens] >pir PH1429 PH1429 Ig heavy chain V region (clone VH5-2R1) - human (fragment) {SUB 1-98} Length = 12
HCRMW16R	HWL0035R	HWLVL77R	нсувн84к	HBJMG15R	н2СВН29R	HWLMT35R
2208	2209	2210	2211	2212	2213	2214

WO 01/22920

pBluescript	pSport1	pBluescript SK-	pBluescript SK-	pCMVSport 3.0	Lambda ZAP II	pCMVSport 2.0	pSport1	pSport1
HKCSJ59	HWMBG21	H2LBB21	H2LAT69	HLWCJ40	нсоск44	НОGDQ <i>57</i>	НWLQM12	HWLVX04
06		100	77	100	62	88	98	98
06		100	64	96	55	85	80	98
328	195	406	393	214	387	314	570	260
71	287	2	1	83	1	3	1	3
6492	6493	6494	6495	6496	6497	6498	6499	0059
gb AAC78645.		gb AAD22960. 1 AF1154	dbj BAA88116 .1	gb AAD17294.	gb AAD28244. 1 AF1242	gb AAD28244. 1 AF1242	dbj BAA33957  .1	gb AAD39916. 1 AF1250
(AF105424) brush border myosin I [Homo sapiens] >gb AAD31189.1 AF127026_1 (AF127026) brush border myosin I [Homo sapiens] >sp AAC78645 AAC78645 Brush border myosin I. >sp AAD31189 AAD31189 Brush border myosin I. >gb AAA20900.1  myosin [Homo sapiens] {SUB		(AF115402) Ets transcription factor ESE-2a [Homo sapiens] >sp AAD22960 AAD22960 Ets transcription factor ESE-2a. >gb AAD22961.1 AF115403_1 (AF115403) Ets transcription factor ESE-2b [Homo sapiens] {SUB 11-265} Length = 265	(AB015335) HRIHFB2072 [Homo sapiens]   >sp[BAA88116[BAA88116 HRIHFB2072 protein (fragment). >sp[AAD26690.1 AF115778_1 (AF115778) short coiled coil protein SCOCO [Mus musculus] {SUB 44-125} Length = 125	(AF120265) tetraspan NET-6 [Homo sapiens] >gb AAD43023.1  (AF100759) transmembrane 4 superfamily protein [Homo sapiens] >sp O95857 O95857 TETRASPAN NET-6. Length = 204	(AF124249) SH2-containing protein Nsp1 [Homo sapiens] >sp Q9Y2X4 Q9Y2X4 SH2-CONTAINING PROTEIN NSP1. Length = 576	taining protein Nsp1 [Homo Q9Y2X4 SH2-CONTAINING Igth = 576		HWLVX04R (AF125099) HSPC038 protein [Homo sapiens] >sp Q9Y5V0 Q9Y5V0 HSPC038 PROTEIN. Length = 76
HKCSJ59R	HWMBG21R	H2LBB21R	H2LAT69R	HLWCJ40R	HCQCK44R	НОGDQ57R	HWLQM12R	HWLVX04R
2215	2216	2217	2218	2219	2220	2221	2222	2223

pBluescript SK-	pSport1	pBluescript SK-	pCMVSport 3.0	pSport1	pSport1	pSport1	pSport1.	pSport1	Uni-ZAP XR	pCMVSport 3.0
H2CBG89	нмгмое8	HCYBM79	HMUB053	HWLOT13	HWLVN81	HWLRV71	HCRMV30	HCROK15	HTOAF87   Uni-ZAP XR	HDPMJ48
59	100	100	93		94	95		97	69	99
38	100	100	92		94	95		26	57	56
317	276	492	518	328	554	137	218	106	345	467
6	-	52	108	2	156	3	3	2	1	81
6501	6502	6503	6504	6505	9059	6507	8059	6209	6510	6511
gb AAD20949.	gb AAD38506. 1 AF1267	gb AAD38133. 1 AF1277	gb AAD17375. 1		gb AAD18080. 1 AAD180	emb CAB5301 5.1		gb AAD44714. 1	emb CAA7141 4.2	gb AAD30564. 1 AF1467
(AF126736) ubiquitin processing protease [Homo sapiens] >sp Q9Y5T5 UBPG_HUMAN UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 16 (EC 3.1.2.15) (UBIQUITIN THIOLESTERASE 16) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 16) (DEUBIQUITINATING ENZYME 16) (UBIQUITIN PROCESSIN	(AF126743) DNAJ domain-containing protein MCJ [Homo sapiens] >sp Q9Y5T4 Q9Y5T4 DNAJ DOMAIN-CONTAINING PROTEIN MCJ. Length = 150	(AF127763) mitogenic oxidase [Homo sapiens] >splQ9Y5S8 Q9Y5S8 MITOGENIC OXIDASE. Length = 564	(AF129075) T-COMPLEX PROTEIN 1, THETA SUBUNIT (TCP-1-THETA) [Homo sapiens] >emb CAA85520.1  Cctq [Homo sapiens] {SUB 33-65} Length = 548		HWLVN81R (AF129756) G5c [Homo sapiens] >sp 095871 095871 NG33. Length = 148	(AJ132584) HIRA-interacting protein HIRIP5 [Homo sapiens] >sp CAB53015 CAB53015 HIRA-interacting protein HIRIP5. Length = 196		(AF138302) decorin variant C [Homo sapiens] >sp Q9Y5N9 Q9Y5N9 DECORIN VARIANT C. Length = 212	Graf protein [Homo sapiens] >sp CAA71414 CAA71414 Graf protein. Length = 759	(AF146793) protein B [Mus musculus] >sp Q9WUP3 Q9WUP3 PROTEIN B (FRAGMENT). Length = 193
H2CBG89R	HWLWQ68R	HCYBM79R	HMUBO53R A	HWLOT13R	HWLVN81R	HWLRV71R	HCRMV30R	HCROK15R	HTOAF87Ra	HDPMJ48R
2224	2225	2226	2227	2228	2229	2230	2231	2232	2233	2234

pSport1	pCMVSport 2.0	Lambda ZAP II	pSport1	pSport1	pSport1	pBluescript SK-	pBluescript SK-	pBluescript SK-	Lambda ZAP II				
HWLNJ72	HOFME52	нсори46	HCRMGSS	HCRNZ49	HASMB62	H2LAD43	H2LAY87	H2LAZ41	НСОСА60	НСQСВ53	нсосн45	нсосл70	нсосг32
97	92	75	92		92		96						
76	99	74	92		84		96						
368	293	298	417	143	121	108	121	178	199	125	229	109	66
39	69	68	52	3	264	16	2	2	2	3	26	2	-
6512	6513	6514	6515	6516	6517	6518	6219	6520	6521	. 6522	6523	6524	6525
gb AAD34057. 1 AF1518	gb AAD34089. 1 AF1518	gb AAD34145. 1 AF1519	gb AAD42874. 1		emb CAB5104		\$6Y5 \$6Y5						
(AF151820) CGI-62 protein [Homo sapiens] >sp Q9Y372 Q9Y372 CGI-62 PROTEIN. Length = 325	(AF151852) CGI-94 protein [Homo sapiens] >sp Q9Y3A2 Q9Y3A2 CGI-94 PROTEIN. Length = 253	(AF151908) CGI-150 protein [Homo sapiens] >sp Q9Y3E8 Q9Y3E8 CGI-150 PROTEIN. Length = 504	HCRMG55R (AF155108) NY-REN-41 antigen [Homo sapiens] >sp Q9Y599 Q9Y599 NY-REN-41 ANTIGEN (FRAGMENT). Length = 241		(AJ243883) putative transcription factor [Periplaneta emb CAB5104 americana] >sp Q9Y071 Q9Y071 PUTATIVE 1.1  TRANSCRIPTION FACTOR. Length = 333		IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414						
HWLNJ72R	HOFME52R	HCQDU46R	HCRMG55R	HCRNZ49R	HASMB62R	H2LAD43R	H2LAY87R	H2LAZ41R	HCQCA60R	нсосвязя	HCQCH45R	HCQCJ70R	HCQCL32R
2235	2236	2237	2238	2239	2240	2241	2242	2243	2244	2245	2246	2247	2248

HCQCP47   Lambda ZAP	HCQDC76 Lambda ZAP	Lambda ZAP II	Lambda ZAP II	HCQDK24 Lambda ZAP	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II	HCQDM76 Lambda ZAP	Lambda ZAP II	HCQDR62 Lambda ZAP	HCQDV85 Lambda ZAP	HCQDW29 Lambda ZAP	HCQDW44 Lambda ZAP II	pSport1
НСОСР47	НСОДС76	нсорн59	нсор182	нсорк24	нсоркз	нсорг42	нсорг82	нсорм76	нсорр62	HCQDR62	нсору85	нсорм29	НСQDW44	HCRPO09
													91	
													82	
163	208	110	245	101	110	186	154	125	110	120	195	242	144	357
2	2	3	42	m	es.	4	∞	3	e.	-	10	3	-	37
6526	6527	6528	6529	6530	6531	6532	6533	6534	6535	6536	6537	6538	6539	6540
													8p Q9Y6Y5 Q9 Y6Y5	
													HCQDW44R IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	
HCQCP47R	HCQDC76R	нсорн59к	HCQDI82R	HCQDK24R	нсоркъзк	HCQDL42R	HCQDL82R	HCQDM76R	<b>НСQDP62R</b>	HCQDR62R	HCQDV85R	HCQDW29R	HCQDW44R	HCRPO09R
2249	2250	2251	2252	2253	2254	2255	2256	2257	2258	2259	2260	2261	2262	2263

pBluescript SK-	pSport1	HKCAA76 Uni-ZAP XR	pSport1	HCQDA64 Lambda ZAP	pSport1	HCQDC26 Lambda ZAP	pSport1	Lambda ZAP II	Lambda ZAP II
HCYBO90	HWLVA95	HKCAA76	HWMCK11	нсорае4	HCRNF45	нсорс26	HWLWN07	нсория	HCQAI55
84		87	85	65	63	81	88	99	
84		85	8	62	61	89	82	62	
204	373	498	429	473	347	269	183	622	333
	44	-	-	3	141	3		89	217
6541	6542	6543	6544	6545	6546	6547	6548	6549	6550
sp Q9Y6Y5 Q9 Y6Y5		emb CAA0918 1.1	emb CAA0918 1.1	emb CAA1156 0.1	emb CAB4685 3.1	emb CAA1667	emb CAA6201 3.1	emb CAA5183	
HCYBO90R   IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414		(AJ010442) immunoglobulin kappa light chain [Homo sapiens] Length = 236	HWMCK11R (AJ010442) immunoglobulin kappa light chain [Homo sapiens] Length = 236	HCQDA64R (AJ223814) striatin [Homo sapiens] >sp 043815 043815 STRIATIN. Length = 780	(AJ388554) hypothetical protein [Canis familiaris] >sp Q9XSR4 Q9XSR4 HYPOTHETICAL 17.5 KD PROTEIN (FRAGMENT). Length = 174	(AL021683) unnamed protein product [Homo sapiens] >sp[O43819 O43819 HYPOTHETICAL 29.8 KD PROTEIN ON CHROMOSOME 22. Length = 266	NBK [Homo sapiens] >emb CAA18260.2  (AL022237) bK1191B2.2 (BCL2-interacting killer (apoptosis-inducing) (NBK, BP4, BIP1)) [Homo sapiens] >gb AAC50413.1  Bik [Homo sapiens] >gb AAC79124.1  apoptosis inducer Nbk [Homo sapiens] >spiens] >gb AAF01156.1  (AF174424) BCL	ribosomal protein L3 [Homo sapiens]  >emb CAA18450.1  (AL022326) dJ333H23.1 (60S Ribosomal Protein L3) [Homo sapiens]  >pir S34195 S34195 ribosomal protein L3, cytosolic  - human >sp P39023 RL3_HUMAN 60S RIBOSOMAL PROTEIN L3 (HIV-1 TAR RNA BINDING PROTEIN B)	
HCYBO90R	HWLVA95R	HKCAA76R	HWMCK11R	HCQDA64R	HCRNF45R	нсорс26R	HWLWN07R	нсориззк	HCQAI55R
2264	2265	2266	2267	2268	2269	2270	2271	2272	2273

pSport1	HCQCK51 Lambda ZAP	Uni-ZAP XR	Uni-ZAP XR	pSport1	pSport1	pSport1
HWLNK89	HCQCK51	HAIDT47	HSUAK69	HCROB90	HCRNI50	HCRPJ34
84	09	88		61	94	83
84	40	82		50	91	80
66	353	53	254	129	577	355
_	108	442	18	335	44	149
6551	6552	6553	6554	6555	9259	6557
emb CAB4322 0.1	emb CAB4326	emb CAB4337		dbj BAA29286	emb CAB5527 6.1	gb AAA59066. 1
	HCQCK51R (AL050097) hypothetical protein [Homo sapiens] >emb CAB43269.1  (AL050097) hypothetical protein [Homo sapiens] >pir T08746/1708746 hypothetical protein DKFZp586B0319.1 - human (fragment) >sp Q9Y3V9 Q9Y3V9 HYPOTHETICAL 17.3 KD PROTEIN (FRAGMENT). Length = 1	(AL050273) hypothetical protein [Homo sapiens] >emb CAB43374.1  (AL050273) hypothetical protein [Homo sapiens] >pir T08720 T08720 ribosomal protein L36 - human >sp Q9Y3U8 Q9Y3U8 60S RIBOSOMAL PROTEIN L36. Length = 105		(AP000001) 106aa long hypothetical protein [Pyrococcus horikoshii] >pir G71244 G71244 hypothetical protein PH0217 - Pyrococcus horikoshii >sp O57956 O57956 HYPOTHETICAL 11.5 KD PROTEIN PH0217. Length = 106	(AL035461) dJ967N21.5 (novel MCM2/3/5 family member) [Homo sapiens] >splCAB55276 CAB55276 6.1 DJ967N21.5 (novel MCM2/3/5 family member) (fragment). Length = 606	18 kDa Alu RNA binding protein [Homo sapiens] >pir A56062 A56062 Alu RNA-binding protein - human Length = 136
HWLNK89R	HCQCK51R	HAIDT47R	HSUAK69RA	HCROB90R	HCRNI50R	HCRPJ34R
2274	2275	2276	2277	2278	2279	2280

HCQBL95 Lambda ZAP	pSport1	pBluescript	pSport1	pSport1	Uni-ZAP XR	pBluescript	pSport1	HCQCO29 Lambda ZAP	pSport1	pSport1	pSport1
нсов 195	HWLOR95	HKCSI32	нwLQК90	HCRNO08	HBGBT36	HKLSA81	HBCJN86	нсосо29	HWLMZ47	HCRNZ75	HCRPD88
	81	76	16	98	85	58	26	95		96	86
	80	76	91	84	84	99	26	93		96	86
195	536	260	496	540	195	345	212	313	282	334	271
124	3	3	2	<b>,</b>	-	112	39	2	82	2	2
6558	6559	0959	6561	6562	6563	6564	9299	9959	6567	8959	6959
	gb AAC52114.	gb AAA36044.	gb AAA36044. 1	gb AAC50495. 1	gb AAA69923.	sp Q29375 RL 7A_PIG	sp Q29214 RL A0_PIG	gb AAB66488. 1		emb CAA2907 5.1	gb AAA52002. 1
	HWLOR95R 3-hydroxyisobutyryl-coenzyme A hydrolase [Homo sapiens] >splQ92931 Q92931 3-HYDROXYISOBUTYRYL-COENZYME A HYDROLASE. Length = 381	40-kDa keratin protein [Homo sapiens] >pir A31370 KRHU9 keratin 19, type I, cytoskeletal - human Length = 400	40-kDa keratin protein [Homo sapiens] >pir A31370 KRHU9 keratin 19, type I, cytoskeletal - human Length = 400	ein kinase, gamma-1 subunit 14619 AAKG_HUMAN S"- ROTEIN KINASE, GAMMA 3AMMA-1 CHAIN). Length	60S ribosomal protein [Mus musculus] >sp P47963 RL13_MOUSE 60S RIBOSOMAL PROTEIN L13 (A52). {SUB 2-213} Length = 213	60S RIBOSOMAL PROTEIN L7A (SURFEIT LOCUS PROTEIN 3) (FRAGMENT). Length = 132	60S ACIDIC RIBOSOMAL PROTEIN P0 (L10E) (FRAGMENT). Length = 93	HCQCO29R (AF014364) beta actin [Cricetinae gen. sp.] >sp 035248 035248 BETA ACTIN. Length = 295		alpha-1 chain precursor (AA -27 to 917) (2953 is 2nd emb CAA2907 base in codon) [Homo sapiens] Length = 944 5.1	alpha-1 type III collagen [Homo sapiens] Length = 345
нсовгэзк	HWLOR95R	HKCSI32R	HWLQK90R	HCRNO08R	HBGBT36R	HKLSA81R	HBCJN86R	HCQCO29R	HWLMZ47R	HCRNZ75R	HCRPD88R
2281	2282	2283	2284	2285	2286	2287	2288	2289	2290	2291	2292

pSport1	HCQDC47 Lambda ZAP	HCQDT07 Lambda ZAP	pBluescript SK-	pSport1	pSport1	pSport1	pSport1
HCRPK90	нсорс47	нсорт07	H2CBR33	HWLXV36	HWLRE24	HWMBA27	HWMBK08
73	100	88	100	96	86	79	97
73	100	84	001	95	86	79	97
235	459	312	242	380	436	228	118
2	y-out	55	3	3	2	-	2
6570	6571	6572	6573	6574	6575	6576	6577
gb AAA52482.	gb AAA36351. 1	dbj BAA90953 .1	emb CAA7618 5.1	emb CAB6305	gb AAA58455. 1	emb CAA8830 7.1	gb AAB08753.
HCRPK90R   alpha-L fucosidase [Homo sapiens] Length = 353	HCQDC47R alpha-N-acetylgalactosaminidase [Homo sapiens] >gb AAA51677.1  alpha-N-acetylgalactosaminidase [Homo sapiens] >gb AAB06718.1  alpha-N-acetylgalactosaminidase [Homo sapiens] >emb CAB41237.1  bK250D10.5 (alpha-N-acetylgalactosaminidase) [Homo sapiens]	HCQDT07R (AK000113) unnamed protein product [Homo sapiens] Length = 273	alternatively spliced form [Homo sapiens] >sp O75666 O75666 CXORF5 (71-7A) PROTEIN (71-7A). Length = 367	HWLXV36R dJ408N23.5 (novel protein simialr to aminopeptidase emb CAB6305   P) [Homo sapiens] >sp CAB63053 CAB63053   DJ408N23.5 (novel protein simialr to aminopeptidase P) (fragment). Length = 135	amplaxin [Homo sapiens] >pir A48063 A48063 manmary tumor/squamous cell carcinoma-associated protein EMS1 - human >sp Q14247 SRC8_HUMAN SRC SUBSTRATE CORTACTIN (AMPLAXIN) (EMS1 ONCOGENE). Length = 550	HWMBA27R aponucin [Homo sapiens] >pirlA57534 A57534 mucin (clone L31) - human (fragment) >sp Q13792 Q13792 APOMUCIN (FRAGMENT). Length = 1042	HWMBK08R arginine-rich protein [Homo sapiens] >pir S27956 S27956 arginine-rich protein - human >sp P55145 ARGR_HUMAN ARGININE-RICH PROTEIN. Length = 234
HCRPK90R	HCQDC47R	HCQDT07R	H2CBR33R	HWLXV36R	HWLRE24R	HWMBA27R	HWMBK08R
2293	2294	2295	2296	2297	2298	2299	2300

pBluescript	pBluescript	pSport1	pSport1	HCQCT96 Lambda ZAP	HCQCV26 Lambda ZAP
HKCSA84	HKCTB10	HCRPX81	HWLRP68	нсост96	HCQCV26
73	75	84	69	77	69
73	92	84	69	73	99
231	102	442	231	176	716
-	434	50	-	m	363
6578	6229	0859	6581	6582	6583
emb CAA5855 6.1	gb AAA24165. 1	gb AAA35762. 1	dbj BAA23322 .1	emb CAA2403	emb CAA2403 1.1
HKCSA84R ARSE [Homo sapiens] >pir I37187 I37187 arylsulfatase E (EC 3.1.6) - human >sp P51690 ARSE_HUMAN ARYLSULFATASE E PRECURSOR (EC 3.1.6) (ASE). Length = 589	aspartokinase II-homoserine dehydrogenase II [Escherichia coli] Length = 810	sociated protein 19618 probable otein - human TELANGIECTASIA TEIN. Length = 588	HWLRP68R ATP synthase subunit e [Homo sapiens] >sp P56385 ATPJ HUMAN ATP SYNTHASE E CHAIN, MITOCHONDRIAL (EC 3.6.1.34). {SUB 2-69} Length = 69	ATPase 6 [Homo sapiens] >gb AAB58948.1  ATPase 6 [Homo sapiens] >pir A01049 PWHU6 H+- transporting ATP synthase (EC 3.6.1.34) protein 6 - human mitochondrion >sp P00846 ATP6_HUMAN ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6). Length = 226	HCQCV26R ATPase 6 [Homo sapiens] >gb AAB58948.1  ATPase 6 [Homo sapiens] >pir A01049 PWHU6 H+- transporting ATP synthase (EC 3.6.1.34) protein 6 - human mitochondrion >sp P00846 ATP6 HUMAN ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6). Length = 226
HKCSA84R	HKCTB10R	HCRPX81R	HWLRP68R	HCQCT96R	HCQCV26R
2301	2302	2303	2304	2305	2306

pSport1	Lambda ZAP II	Uni-ZAP XR	pSport1	pBluescript SK-	pSportl	pSport1	pSport1	pSport1	pSport1	pSport1
HWLXR95	HCQCV96 Lambda ZAP	HEPAD45 Uni-ZAP XR	HCRNP41	HCYBK83	HCRND59	HCRNE86	HWLQZ23	HCRMA15	HCRMJ42	HCRMK50
100	61	100	63		86					100
100	58	100	09		96					100
253	368	277	281	01	385	117	231	152	244	113
2	147	2	39	471	2	-	-	9	14	24
6584	6585	6586	6587	6588	6589	0659	6591	6592	6593	6594
gb AAA51807.	dbj BAA07295 .1	gb AAB51444. 1	dbj BAA35182 .1		gb AAA30384. 1					emb CAB6919 5.1
ATPase coupling factor 6 subunit [Homo sapiens] >emb CAB53667.1  (AL110183) hypothetical protein [Homo sapiens] >emb CAB53667.1  (AL110183) hypothetical protein [Homo sapiens] >pir JT0563 JT0563 coupling factor 6 precursor, mitochondrial - human >sp P1885	ATPase subunit 6 [Homo sapiens] >dbj BAA07295.1  ATPase subunit 6 [Homo sapiens] Length = 226	autoantigen [Homo sapiens] >sp Q13025 Q13025 AUTOANTIGEN. Length = 968	(AB021288) beta 2-microglobulin [Homo sapiens] >gb AAA87972.1  beta-2-microglobulin [Pan troglodytes] >gb AAA88008.1  beta-2-microglobulin [Gorilla gorilla] >gb AAA88008.1  (AF072097) beta-2 microglobulin [Homo sapiens] >pir A90976 MGHUB2 beta-2-microglob		beta-adrenergic receptor kinase [Bos taurus] >pir A40088 A40088 beta-adrenergic-receptor kinase (EC 2.7.1.126) 1 - bovine >sp P21146 ARK1 BOVIN BETA-ADRENERGIC RECEPTOR KINASE 1 (EC 2.7.1.126) (BETA-ARK-1) (G- PROTEIN COUPLED RECEPTOR KINASE 2). Length =					unnamed protein product [unidentified] Length = 180
HWLXR95R	HCQCV96R	HEPAD45R	HCRNP41R	HCYBK83R	HCRND59R	HCRNE86R	HWLQZ23R	HCRMA15R	HCRMJ42R	HCRMK50R
2307	2308	2309	2310	2311	2312	2313	2314	2315	2316	2317

pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1										
HCRMO88	HCRNB87	HCRNI95	HCRNL44	HCROD06	HCROT10	HCRPF12	HCRPK22	HCRPK46	HCRPK48	HCRQF74	HCRQG02	HCRQM26	нском90	HHMMA34	HHMMA44	HHIMMC42	HHMMC86	HHMMD59	HHMME38	HHMME40	HHMME50	HHMME58	HHMME80	HHMMF60	HHMMF79	HOCTA39	HS2AN66	HULCG37
										100	100		100								100							
										100	95		100								100							
352	995	26	561	454	211	231	96	142	153	129	171	175	104	157	219	160	154	143	245	95	110	142	159	136	173	582	391	166
92	354	2	1	137	2	25	1	29	22		-	23	٣	14	26	26	14	9	111	3	3	5	25	2	3	457	161	44
6595	9659	6597	8659	6299	0099	1099	6602	6603	6604	9099	9099	2099	8099	6099	6610	6611	6612	6613	6614	6615	9199	6617	6618	6619	6620	6621	6622	6623
										unnamed protein product [unidentified] Length = 180 emb CAB6919 5.1	unnamed protein product [unidentified] Length = 180 emb CAB6919 5.1		HCRQM90R unnamed protein product [unidentified] Length = 180 emb CAB6919   5.1								HHMME50R unnamed protein product [unidentified] Length = 180 emb CAB6919   5.1							
HCRMO88R	HCRNB87R	HCRNI95R	HCRNL44R	HCROD06R	HCROT10R	HCRPF12R	HCRPK22R	HCRPK46R	HCRPK48R	HCRQF74R	HCRQG02R	HCRQM26R	HCRQM90R	HHMMA34R	HHMMA44R	HHMMC42R	HHMMC86R	HHMMD59R	HHMME38R	HHMME40R	HHMME50R	HHMME58R	HHMME80R	HHMMF60R	HHMMF79R	HOCTA39R	HS2AN66R	HULCG37R
2318	2319	2320	2321	2322	2323	2324	2325	2326	2327	2328	2329	2330	2331	2332	2333	<del>                                     </del>	2335	2336	2337		2339	2340	2341	2342	2343	2344	2345	2346

Ē	Ŧ	11	ד	E	E	£	£	E	£	£	Ę	Ŧ	Ŧ	II.	Ŧ	£	E E	£	E	뒫	ĮĘ.	т Т	Ŧ	<u> </u>	£	핕	듣	E	£
pSport1	pSport1	pSport1	pSportl	pSportl	pSportl	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSportl	pSport1	pSport1	pSport1								
HWLM016	HWLM029	HWLM044	HWLM047	HWLM084	HWLMQ27	HWLMQ28	HWLMQ64	НWLМQ65	HWLNZ01	HWLNZ20	HWLNZ35	HWLNZ44	HWLNZ59	HWLOV04	HWLOW58	HWLOZ37	HWLQF96	HWLRV12	HWLWE68	HWLXA42	HWLXA48	HWLXA94	HWLXI52	HWMBR11	HWMBR26	HWMBR49	HWMBR53	HWMBR56	HWMBS18
HW	HW	HW	HW	HW	HW	HW	HM	HW	H	H	H	HA	HA	70 HW	HW	HA	HV	H	HM	100 HW	HM	HW	H	HW	HW	100 HW	HM	HM	HM
														62 7						100						100			
146	143	142	129	177	184	246	168	197	101	121	158	181	184	134 (	464	513	464	120	241	176 1	132	186	114	136	278	260.   1	194	156	117
42	24	35	31	1	2	19		99	3	2	3	2	2	3	264	322	249	7	2	ы	1	10	1	2	3	48	3	1	1
6624	6625	9799	6627	8799	6629	0630	6631	6632	6633	6634	6635	9636	6637	8638	6639	6640	6641	6642	6643	6644	6645	9499	6647	6648	6649	0599	6651	6652	6653
														emb CAB6919 5.1						180 emb CAB6919 5.1						emb CAB6919   5.1			
A CANADA TANADA														HWLOV04R unnamed protein product [unidentified] Length = 180 emb CAB6919   5.1						unnamed protein product [unidentified] Length =						HWMBR49R   unnamed protein product [unidentified] Length = 180 emb CAB6919   5.1			
HWLM016R	HWLMO29R	HWLMO44R	HWLM047R	HWLMO84R	HWLMQ27R	HWLMQ28R	HWLMQ64R	HWLMQ65R	HWLNZ01R	HWLNZ20R	HWLNZ35R	HWLNZ44R	HWLNZ59R	HWLOV04R	HWLOW58R	HWLOZ37R	HWLQF96R	HWLRV12R	HWLWE68R	HWLXA42R	HWLXA48R	HWLXA94R	HWLXI52R	HWMBR11R	HWMBR26R	HWMBR49R	HWMBR53R	HWMBR56R	HWMBS18R
2347	2348	2349	2350	2351	2352	2353	2354	2355	2356	2357	2358	2359	2360	2361	2362	2363	2364	2365	2366	2367	2368	5369	2370	2371	2372	2373	2374	2375	2376

						1				Γ			_	_		r—			~					10.
pSport1	pSport1	pSportl	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	Uni-ZAP XR	pSport1	pSport1	pSport1	pSport1	Lambda ZAP II									
HWMBS36	HWMBS81	HWMBW16	HCRMH48	HCRMY75	HCRNZ66	HCROJ91	HCRPR59	HCRPY45	HCRQG95	HHMMC37	HHMMC68	HHIMIMD73	HHMMF44	HTWEL13	HWLMT48	HWLQK72	HWLUI68	HWLVY86	HE2JQ95	HCRMH46	HWLMW81	HWLND45	HWLWG95	нсосхія
									98														-	73
									82															59
187	223	394	421	254	305	283	202	430	185	144	137	160	154	157	480	103	476	464	232	334	191	501	298	143
2	5	98	98	3	3	<b>∞</b>	26	218	3	13	3	2	2	2	343	17	3	234	2	65	3	91	23	3
6654	6655	9599	6657	8599	699	0999	1999	6662	6999	6664	9999	9999	2999	8999	6999	0299	6671	6672	6673	6674	6675	9299	2299	8299
									emb CAB6919 5.1															gb AAA52645.
		2							unnamed protein product [unidentified] Length = 180												2			beta-hexosaminidase beta-subunit [Homo sapiens] >pir A31250 A31250 beta-N-acetylhexosaminidase (EC 3.2.1.52) beta chain precursor - human >sp P07686 HEXB_HUMAN BETA-HEXOSAMINIDASE BETA CHAIN PRECURSOR (EC 3.2.1.52) (N-ACETYL-BETA-GLUCOSAMINIDASE) (BETA-N
HWMBS36R	HWMBS81R	HWMBW16R	HCRMH48R	HCRMY75R	HCRNZ66R	HCROJ91R	HCRPR59R	HCRPY45R	HCRQG95R	HHMMC37R	HHMMC68R	HHMMD73R	HHMMF44R	HTWEL13RA	HWLMT48R	HWLQK72R	HWLUI68R	HWLVY86R	HE2JQ95R	HCRMH46R	HWLMW81R	HWLND45R	HWLWG95R	нсосх19к
2377	2378	2379	2380	2381	2382	2383	2384	2385	2386	2387	2388	2389	2390	2391	2392	2393	2394	2395	2396	2397	2398	2399	2400	2401

7	Ŧ	£	Ŧ	Ŧ	Ŧ	딛	=
pSportl	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1
<b>НС</b> РООЗЗ	HCRNT56	HCRPR45	HCRMJ70	HWLMM72	HCRMD32	HCRQC71	HWMCH04
63 H(	PH 67	100 Н	100 H	87 HV	75 H(	86 H	84 HX
59 6	78 7	100	100   10	81 8	73 7	6	8
442	516	167	169	385	292	261	320
2		8	2	08	2	-	<sub>ε</sub>
6299	0899	6681	6682	899	6684	9895	9899
gb AAB51199. 1	gb AAA51838. 1	gb AAB20222.	emb CAA3103 7.1	gb AAB51329.	gb AAC16021.	gb AAA30423.	gb AAA51908. 1
HCRQO33R   bHLH protein MesP2 [Mus musculus]   >sp 008574 008574 MESODERM POSTERIOR 2 (BHLH PROTEIN MESP2). Length = 370	BN51 protein [Homo sapiens] >pir A43700 A43700 BN51 protein - human >sp P05423 BN51_HUMAN BN51 PROTEIN. Length = 395	branched-chain alpha-keto acid dehydrogenase E1 alpha subunit [human, Peptide, 443 aa] [Homo sapiens] Length = 443	[Homo sapiens] 1 snRNP protein C - human JMAN U1 SMALL CLEOPROTEIN C (U1-C).	HWLMM72R C10 [Homo sapiens] >sp Q99622 Q99622 CHROMOSOME 12P13 SEQUENCE; HTGS PHASE 3, COMPLETE SEQUENCE. Length = 126	HCRMD32R CAG-isl 7 [Homo sapiens] >sp P50914 RL14_HUMAN 60S RIBOSOMAL PROTEIN L14 (CAG-ISL 7). {SUB 2-213} Length = 213	calpactin I light chain [Bos taurus] >gb AAA58404.1  gb AAA30423 calpactin I light chain [Homo sapiens] 1  >gb AAA58426.1  cellular ligand of annexin II [Homo sapiens] >pir JC1139 JC1139 calpactin I light chain - human >pir B28489 B28489 calpactin I light chain - bovine	carbonic anhydrase II [Homo sapiens] >gb AAA51909.1  carbonic anhydrase II [Homo sapiens] >gb AAA51911.1  carbonic anhydrase II [Homo sapiens] >emb CAA68426.1  carbonic anhydrase II (AA 1-260) [Homo sapiens] >pir A27175 CRHU2 carbonate dehydratase (EC 4.2)
HCRQ033R	HCRNT56R	HCRPR45R	HCRMJ70R	HWLMM72R	HCRMD32R	HCRQC71R	HWMCH04R
2402	2403	2404	2405	2406	2407	2408	2409

HCQDH40 Lambda ZAP	HKAHM80 pCMVSport 2.0	pBluescript SK-	HCQAN45 Lambda ZAP	pSport1	pSport1	pSport1	pSport1
нсорн40	НКАНМ80	H2CBM60	HCQAN45	HWMCI76	HWLXR73	HWLOI59	HWLUX53
	85	66	69			82	87
	88	66	99			08	87
331	388	532	281	317	341	646	282
2	2	2	E.	3	3	2	_
2899	8899	6899	0699	6691	6695	6693	6694
	gb AAB04798. 1	dbj BAA16470 .1	gb AAA36022. 1			gb AAB02683. 1	emb CAA4928 8.1
	HKAHM80R CDC37 homolog [Homo sapiens] >gb AAB63979.1  gb AAB04798. CDC37 homolog [Homo sapiens]	CDP-DIACYLGLYCEROLSERINE O-PHOSPHATIDYLTRANSFERASE (EC 2.7.8.8) (PHOSPHATIDYLSERINE SYNTHASE). [Escherichia coli] >dbj BAA16473.1  CDP-DIACYLGLYCEROLSERINE O-PHOSPHATIDYLTRANSFERASE (EC 2.7.8.8) (PHOSPHATIDYLSERINE SYNTHASE). [Escherichia coli] >sp P7	chaperonin (HSP60) [Homo sapiens] >sp G242370 G242370 P60, 60-KDA HEAT SHOCK PROTEIN, HSP60. {SUB 27-55} >pir A56868 A56868 heat shock protein 60 - bovine (fragment) {SUB 27-48} Length = 573			HWLOI59R tat interactive protein [Homo sapiens] Length = 482	HWLUX53R cpn10 protein [Bos taurus] >gb AAA50953.1  chaperonin 10 [Homo sapiens] >emb CAA53455.1  heat shock protein 10 [Homo sapiens] >emb CAB75425.1  (AJ250915) chaperonin 10, Hsp10 protein [Homo sapiens] >pir S47532 S47532 chaperonin groES - human >pir A56682 A
HCQDH40R	HKAHM80R	Н2СВМ60К	HCQAN45R	HWMCI76R	HWLXR73R	HWLOI59R	HWLUX53R
2410	2411	2412	2413	2414	2415	2416	2417

pSport1	pCMVSport 3.0	HCQCQ58 Lambda ZAP	НСQDQ11 Lambda ZAP II	HCQDM81 Lambda ZAP
HCRNB29	HARMO20	нсосозя	нсороп	нсорм81
100	64	9	75	72
100	09	64	72	71
336	275	280	550	251
-	153	158	272	129
6695	9699	7699	8699	6699
gb AAB31350. 1	emb CAA2403	dbj BAA77671	dbj BAA77671	emb CAA2403
cyclophilin C, Cyp-C [human, kidney, Peptide, 212 aa] [Homo sapiens] >pir A54204 A54204 peptidylprolyl isomerase (EC 5.2.1.8) C precursor - human >sp P45877 CYPC_HUMAN PEPTIDYL-PROLYL CIS-TRANS ISOMERASE C (EC 5.2.1.8) (PPIASE) (ROTAMASE) (CYCLOPHILIN C).	cytochrome oxidase III [Homo sapiens] >pir A00482 OTHU3 cytochrome-c oxidase (EC 1.9.3.1) chain III - human mitochondrion >gb AAB63452.1  (AF004341) cytochrome c oxidase subunit III [Homo sapiens] {SUB 167-261} Length =	cytochrome c oxidase subunit 3 [Homo sapiens] >dbj BAA77671.1  cytochrome c oxidase subunit 3 [Homo sapiens] >sp P00414 COX3_HUMAN CYTOCHROME C OXIDASE POLYPEPTIDE III (EC 1.9.3.1). >sp BAA77671 BAA77671 Cytochrome c oxidase subunit 3 (fragment).	cytochrome c oxidase subunit 3 [Homo sapiens] >dbj BAA77671.1  cytochrome c oxidase subunit 3 [Homo sapiens] >sp P00414 COX3_HUMAN CYTOCHROME C OXIDASE POLYPEPTIDE III (EC 1.9.3.1). >sp BAA77671 BAA77671 Cytochrome c oxidase subunit 3 (fragment).	HCQDM81R cytochrome oxidase III [Homo sapiens]  >pir A00482 OTHU3 cytochrome-c oxidase (EC 1.9.3.1) chain III - human mitochondrion  >gb AAB63452.1  (AF004341) cytochrome c oxidase subunit III [Homo sapiens] {SUB 167-261} Length = 261
HCRNB29R	HARMO20R	HCQCQ58R	нсороня	HCQDM81R
2418	2419	2420	2421	2422

HCQCU09 Lambda ZAP	HHSGT23 Uni-ZAP XR	Lambda ZAP II	pSport1	Uni-ZAP XR
нсосио9	HHSGT23	нсорап	HFIJB15	HACCH14
68	55	62	87	67
83	51	59	84	63
186	284	284	517	267
70	165	165	326	100
0400	6701	6702	6703	6704
emb CAA6763 0.1	gb AAA20843.	gb AAA20843.	gb AAA20843. 1	gb AAA20843.
HCQCU09R cytochrome oxidase subunit I [Hylobates lar] >dbj BAA07496.1  Cytochrome C oxidase subunit 1 (COXI) [Hylobates syndactylus] >dbj BAA07496.1  Cytochrome C oxidase subunit 1 (COXI) [Hylobates syndactylus] >pir I37049 I37049 cytochrome-c oxidase (EC 1.9.3.1)	cytochrome oxidase subunit II [Homo sapiens] >gb AAA20844.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA20845.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA20847.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA31850.1  cytochrome	cytochrome oxidase subunit II [Homo sapiens] >gb AAA20844.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA20845.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA20847.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA31850.1  cytochrome	cytochrome oxidase subunit II [Homo sapiens] >gb AAA20844.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA20845.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA20847.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA31850.1  cytochrome	cytochrome oxidase subunit II [Homo sapiens] >gb AAA20844.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA20845.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA20847.1  cytochrome oxidase subunit II [Homo sapiens] >gb AAA31850.1  cytochrome
HCQCU09R	HHSGT23R	HCQDAIIR	HFIJB15R	HACCH14R
2423	2424	2425	2426	2427

pSportl	pSport1	pSport1	pSport1	pSport1	pCMVSport 3.0	pSport1	pSport1
HWLNH49	HWLR011	HCRMP82	HCRPV08	HWMBB77	ннерг48	HCRPT53	HA5BV11
74	98	90	22	92	70	100	09
89	86	06	55	92	70	100	58
401	190	475	569	415	229	179	314
267	2	2	3	2	2	3	3
6705	9029	6707	6708	6109	6710	6711	6712
gb AAA02993.	gb AAB21814.	gb AAA52144. 1	sp Q29386 Q2 9386	9.1	gb AAC51226. 1	emb CAA5853 5.1	emb CAA6185 7.1
HWLNH49R   cytochrome P450 PCN3 [Homo sapiens]	HWLRO11R cytochrome P450 reductase {EC 1.6.2.4} [human, placenta, Peptide Partial, 676 aa] [Homo sapiens] Length = 676	051	KERATIN TYPE II CYTOSKELETAL 8 (FRAGMENT). Length = 116	HWMBB77R macrophage inflammatory protein-2beta precursor [Homo sapiens] >pirJH0282 B38290 GRO-gamma precursor - human >sp P19876 MI2B_HUMAN MACROPHAGE INFLAMMATORY PROTEIN-2-BETA PRECURSOR (MIP2-BETA) (GROWTH REGULATED PROTEIN GAMMA) (GRO-GAMMA) >gb AAA03454.1	cytotoxic ligand TRAIL receptor [Homo sapiens] >sp 000220 000220 CYTOTOXIC LIGAND TRAIL RECEPTOR. Length = 468	DAP-3 [Homo sapiens] >sp P51398 DAP3_HUMAN   emb CAA5853 DEATH-ASSOCIATED PROTEIN 3 (DAP-3). 5.1  Length = 398	death associated protein 5 [Homo sapiens] Length = 907
HWLNH49R	HWLRO11R	HCRMP82R	HCRPV08R	HWMBB77R	HHEPL48R	HCRPT53R	HA5BV11R
2428	2429	2430	2431	2432	2433	2434	2435

PCT/US00/26524

HCQCD92 Lambda ZAP	HTXJU67 Uni-ZAP XR	3 pSport1	6 pSport1	HCQCO67 Lambda ZAP	pSport1	pSport1	5 pSport1	pBluescript SK-
НСОСО92	HTXJU67	HWMCL33	HCRMM56	190000н	HCRPQ41	HWLVI33	HWMBASS	HCYBK79
<i>L</i> 9	06	68		79	65	70	54	
57	06	88		73	45	65	53	
130	124	367	250	209	320	298	404	217
2	23	2	2	m	3	2	٤	=
6713	6714	6715	6716	6717	6718	6719	6720	6721
gb AAA52167.	gb AAC34803.	gb AAA98749. 1		emb CAB1655 6.1	dbj BAA09454 .1	emb CAA8106 6.1	gb AAC50423.	
HCQCD92R   decay-accelerating factor precursor [Homo sapiens]   Length = 376	liacylglycerol kinase alpha [Homo th = 210	HWMCL33R diacylglycerol kinase [Homo sapiens] >sp P52824 KDGD_HUMAN DIACYLGLYCEROL KINASE, DELTA (EC 2.7.1.107) (DIGLYCERIDE KINASE) (DGK- DELTA) (DAG KINASE DELTA) (80 KD DIACYLGLYCEROL KINASE). Length = 942		dJ142L7.3 (Connective tissue growth factor (NOV, GIG) LIKE protein) [Homo sapiens] >gb AAD31517.1 AF143679_1 (AF143679) lost in inflanmatory breast cancer tumor suppressor protein [Homo sapiens] >sp O95958 O95958 DJ142L7.3 (CONNECTIVE TISSUE GROWTH FACTOR	DOCK180 protein [Homo sapiens] db >sp Q14185 Q14185 DOCK180 PROTEIN. Length = .1  1865	dodecenoyl-CoA delta-isomerase [Homo sapiens] >pir A55723 A55723 dodecenoyl-CoA Delta-isomerase (EC 5.3.3.8) precursor, mitochondrial-human >sp P42126 D3D2_HUMAN 3,2-TRANS-ENOYL-COA ISOMERASE, MITOCHONDRIAL PRECURSOR (EC 5.3.3.8) (DODECENOYL-COA DELTA-I	HWMBA55R   dynamitin [Homo sapiens]   >sp Q13561 DYNC_HUMAN DYNACTIN, 50 KD   ISOFORM (50 KD DYNEIN-ASSOCIATED   POLYPEPTIDE) (DYNAMITIN). Length = 406	
нсоср92к	HTXJU67R	HWMCL33R	HCRMM56R	нсособ7к	HCRPQ41R	HWLVI33R	HWMBA55R	HCYBK79R
2436	2437	2438	2439	2440	2441	2442	2443	2444

PCT/US00/26524

2445	HCRPS75R	elongation factor 2 [Homo sapiens]  >emb CAA77750.1  human elongation factor 2 [Homo sapiens] >pir S18294 EFHU2 translation elongation factor eEF-2 - human >sp P13639 EF2_HUMAN ELONGATION FACTOR 2 (EF-2). >gb AAA50388.1  elongation factor 2 [Homo sapiens]	emb CAA3582 9.1	6722	2	328	88	88	HCRPS75	pSport1
2446	HCRMJ60R	endoglin [Homo sapiens] Length = 625	emb CAA5089 1.1	6723	81	230	11	78	HCRMJ60	pSport1
2447	HCRPV09R	endozepine precursor [Homo sapiens]  >pir B26448 NZHU endozepine - human  >sp P07108 ACBP_HUMAN ACYL-COA-BINDING PROTEIN (ACBP) (DIAZEPAM BINDING INHIBITOR) (DBI) (ENDOZEPINE) (EP). {SUB 2-87} Length = 87	gb AAA35788. 1	6724	1	318	100	100	HCRPV09	pSport1
2448	HCRON89R	enhancer protein [Homo sapiens] >pir I54533 I54533 enhancer protein - human Length = 199	gb AAA50464. 1	6725	2	361	08	08	HCRON89	pSport1
2449	HAAAT79R	HAAAT79R epithelial glycoprotein (EGP) precursor [Homo sapiens] Length = 314	gb AAA35723.	6726		579	66	66	HAAAT79	pSport1
2450	HLDDP53R	HLDDP53R ERD-2-like protein, ELP-1 - human pir A42286 A4 >emb CAA45277.1  KDEL receptor [Homo sapiens] 2286 {SUB 3-214} Length = 214	pir A42286 A4 2286	6727	28	486	86	86	HLDDP53	pCMVSport 3.0
2451	HWLME23R	HWLME23R FAST kinase [Homo sapiens] >pir I37386 I37386 FAST kinase - human >sp Q14296 Q14296 FAST KINASE. Length = 549	emb CAA6044 8.1	6728	3	407	53	53	HWLME23	pSport1
2452	HWLVP88R	HWLVP88R fau [Homo sapiens] >emb CAA46714.1  fau 1 [Homo sapiens] >pir JC1278 JC1278 ubiquitin-like protein / ribosomal protein S30, cytosolic - human >sp P35544 UBIM_HUMAN_UBIQUITIN-LIKE PROTEIN FUBI. {SUB 1-74} Length = 133	emb CAA4671 6.1	6729	3	473	77	79	HWLVP88	pSport1

PCT/US00/26524

	•				
pSport1	HMWAM77 Uni-ZAP XR	HCQCO75 Lambda ZAP	Lambda ZAP II	pSport1	pSport1
HCROF57	HMWAM77	нсосо75	HCQCR94	HNBTG35	HCROE12
96	09	52	95	92	
96	57	46	95	91	
493	78	82	114	285	105
2	-	264	_		10
6730	6731	6732	6733	6734	6735
gb AAB49652.	sp Q05932 FO LC_HUMAN	pir I43010 QQ BPGL	emb CAA4839 4.1	gb AAF00111. 1 AF1348	
/ATION	HMWAM77R FOLYLPOLYGLUTAMATE SYNTHASE, MITOCHONDRIAL PRECURSOR (EC 6.3.2.17) (FOLYLPOLY-GAMMA-GLUTAMATE SYNTHETASE) (FPGS). >gb AAA35852.1  folylpolyglutamate synthetase [Homo sapiens] {SUB 43-587} >gb AAA87568.1  folylpolyglutamate synthetase [Homo sapiens] {SUB 1	git protein - phage lambda >sp P03762 SIEB_LAMBD SUPERINFECTION EXCLUSION PROTEIN B. >emb CAA23982.1  reading frame GIT [bacteriophage lambda] {SUB 5-183} >emb CAA41524.1  git [bacteriophage lambda] {SUB 6-67} Length = 183	11_	HNBTG35R (AF134895) glyoxylate reductase [Homo sapiens]  >gb AAD45886.1 AF146018_1 (AF146018)  hydroxypyruvate reductase [Homo sapiens]  >gb AAD46517.1 AF146689_1 (AF146689)  hydroxypyruvate reductase [Homo sapiens]  >sp AAD45886 AAD45886 Hydroxypyruvate reductase. >sp	
HCROF57R	HMWAM77R	HCQCO75R	HCQCR94R	HNBTG35R	HCROE12R
2453	2454	2455	2456	2457	2458

Lambda ZAP II	Lambda ZAP II	HCQAM84 Lambda ZAP	pSport1	HCQCF55 Lambda ZAP	pSportl	pSport1
нсолве	HCQAR52	HCQAM84	HWLMG29		HCRMV87	HWLWB88
98	83	70	100	100	88	
84	82	63	86	100	83	
408	103	131	204	274	328	108
118	2	184		2	260	-
6736	6737	6738	6739	6740	6741	6742
gb AAA35933.	gb AAA35933.	gb AAA18898. 1	emb CAA3680 3.1	emb CAA2713 7.1	sp Q29222 Q2 9222	
gro protein [Homo sapiens] >emb CAA31027.1  MGSA preprotein (AA -34 to 73) [Homo sapiens] >emb CAA38361.1  melanoma growth stimulatory activity preprotein [Homo sapiens] >pir S13669 A28414 melanoma growth-stimulatory activity precursor - human >sp P09341	gro protein [Homo sapiens] >emb CAA31027.1  MGSA preprotein (AA -34 to 73) [Homo sapiens] >emb CAA38361.1  melanoma growth stimulatory activity preprotein [Homo sapiens] >pir S13669 A28414 melanoma growth-stimulatory activity precursor - human >sp P09341	growth-regulating protein [Homo sapiens] >pir A56008 A56008 growth-regulating protein BB1 - human >sp P50290 BB1_HUMAN GROWTH- REGULATING PROTEIN BB1. Length = 57	HWLMG29R GTP binding protein [Mus musculus]  >pir A39611 A39611 probable GTP-binding protein - 3.1 mouse >sp P23249 MV10_MOUSE PROTEIN MOV-10. >emb CAA53453.1  gb 110 /Mov 10 locus [Mus musculus] {SUB 1-45} Length = 1004	alpha subunit (aa 1-394) [Bos taurus] >sp P04896 GBAS_BOVIN GUANINE NUCLEOTIDE-BINDING PROTEIN G(S), ALPHA SUBUNIT (ADENYLATE CYCLASE- STIMULATING G ALPHA PROTEIN). Length = 394	HCRMV87R GTP_BINDING PROTEIN (FRAGMENT). Length = 92	
HCQAB69R	HCQAR52R	HCQAM84R	HWLMG29R	HCQCF55R	HCRMV87R	HWLWB88R
2459	2460	2461	2462	2463	2464	2465

2466	HODGF21R	HODGF21R (AF028832) Hsp89-alpha-delta-N [Homo sapiens] >sp O75322 O75322 HSP89-ALPHA-DELTA-N. Length = 539	gb AAC25497. 1	6743	٣	242	94	46	HODGF21	HODGF21 Uni-ZAP XR
2467	HWLMC42R	HWLMC42R non-histone protein HMG2 precursor [Sus scrofa] >pir <sub>1</sub> A34719 A34719 nonhistone chromosomal protein HMG-2 - pig >sp <sub>1</sub> P17741 HMG2_PIG HIGH MOBILITY GROUP PROTEIN HMG2 (HMG-2). {SUB 2-210} Length = 210	gb AAA31051. 1	6744	7	255	08	93 1	HWLMC42	pSport1
2468	HCRQJ58R	histamine H1 receptor [Homo sapiens]  >dbj BAA03319.1  histamine H1 receptor [Homo sapiens] >emb CAA54182.1  histamine H1 receptor [Homo sapiens] >emb CAA84380.1  Human histamine H1 receptor [Homo sapiens]  >gb AAB95156.1  (AF026261) histamine H1 receptor [	dbj BAA05840 .1	6745	29	478	93	93	HCRQJ58	pSport1
2469	HCRQC27R	histone H2A variant (AA 1-141) [Drosophila melanogaster] >emb CAA33555.1  histone H2A [Drosophila melanogaster] >pir S08118 S08118 histone H2A.vD - fruit fly (Drosophila melanogaster) >sp P08985 H2AV_DROME HISTONE H2A VARIANT. {SUB 2-141} >gb AAA72378.1	emb CAA3037 0.1	6746	135	224	81	68	HCRQC27	pSport1
2470	HWLXR58R	HWLXR58R HMG-1 [Homo sapiens] >sp Q14321 Q14321 HMG- dbj BAA09924   1. Length = 215	dbj BAA09924 .1	6747	1	384	100	100	HWLXR58	pSport1
2471	HMWHX32R	HMWHX32R [HsMcm6 [Homo sapiens] >sp[Q14566]MCM6_HUMAN DNA REPLICATION LICENSING FACTOR MCM6 (P105MCM). Length = 821	dbj BAA12699 .1	6748	1	180	100	100 H	HMWHX32	100 HMWHX32 Uni-ZAP XR
2472	HCROW95R	HCROW95R   HTF4a 5"-region hypothetical 13K protein - human   Length = 117	pir A56611 A5 6611	6749	317	3	83	84	HCROW95	pSport1
2473	HCYBO60R	HCYBO60R   HU-K4 [Homo sapiens] >sp Q92853 Q92853 HU-   K4. Length = 437	gb AAB16799. 1	6750	25	387	74	9/	HCYBO60	pBluescript SK-

HE2BG62 Uni-ZAP XR	W12 pSport1	K90 Uni-ZAP XR	F23 pCMVSport 1	F61 pSport1	R28 pBluescript	P47 pSport1	G58 pSport1
HE2BC	HCRMW12	нкрек90	HHBEF23	HWLVF61	HKCSR28	HWMBP47	HWMBG58
94	92	62	52	96	100	8	91
88	92	57	47	8	100	84	91
336	496	188	269	2	190	171	333
187	2	18	186	382	∞	-	446
6751	6752	6753	6754	6755	6756	6757	6758
gb AAC05579.	dbj BAA02656 .1	dbj BAA12865 .1	pirJC1348JC1 348	sp Q16465 YZ A1_HUMAN	gb AAB40244. 1	emb CAA1667 0.1	gb AAD05167.
hunnan gamma-glutamyl hydrolase [Homo sapiens] >gb AAF03360.1  (AF147083) gamma-glutamyl hydrolase [Homo sapiens] >sp Q92820 GGH_HUMAN GAMMA-GLUTAMYL HYDROLASE PRECURSOR (EC 3.4.19.9) (GAMMA-GLU-X CA\ROXYPEPTIDASE) (CONJUGASE) (GH). >sp AAF03360 AAF03360 G	HCRMW12R human homolog of DnaJ protein [Homo sapiens] Length = 397	golo	hypothetical 18K protein - goldfish mitochondrion Length = 166	HWLVF61R HYPOTHETICAL PROTEIN (FRAGMENT). Length = 122	HKCSR28R hypothetical protein [Escherichia coli] >gb AAC73592.1  (AE000155) putative ATP- binding component of a transport system [Escherichia coli] >pir A64780 A64780 probable ABC ransport protein ybbL - Escherichia coli >sp P77279 YBBL_ECOLI HYPOTHETICAL ABC TRAN	HWMBP47R (AL021682) unnamed protein product [Homo emb CAA1667 sapiens] >sp O43788 O43788 CDNA MAPPING TO  0.1  22Q13. Length = 287	HWMBG58R (AF106966) I3 protein [Homo sapiens] >sp O95415 O95415 I3 PROTEIN. Length = 125
HE2BG62R	HCRMW12R	HRDEK90R	HHBEF23R	HWLVF61R	HKCSR28R	HWMBP47R	HWMBG58R
2474	2475	2476	2477	2478	2479	2480	2481

pSport1	pSport	pSport1	pSport1	pSport1	pSport1	HCQAV48 Lambda ZAP II	pSport1
HCRMF92	HWLQF89	HWLOG90	HWLRV24	HWMCC54	HWLNK85	HCQAV48	HWMCM79
74	92	79	81	94	98	78	62
<i>L</i> 9	92	75	81	06	8	78	55
368	263	501	241	395	208	243	209
240	en en	49	2	es.	2	-	78
6759	0929	6761	6762	6763	6764	6765	9929
emb CAA5136	emb CAA5136 0.1	gb AAD30821. 1	pir JE0241 JE0 241	emb CAA5113	gb AAF21612. 1	emb CAA8169	gb AAA02610. 1
IEF 7442 [Homo sapiens] >gb AAC50231.1  retinoblastoma-binding protein RbAp46 [Homo sapiens] >gb AAC36349.1  (AF090306) retinoblastoma binding protein [Rattus norvegicus] >pir 139181 139181 G1/S transition control protein-binding protein RbAp46 - human >s	HWLQF89R IEF 7442 [Homo sapiens] >gb AAC50231.1  retinoblastoma-binding protein RbAp46 [Homo sapiens] >gb AAC36349.1  (AF090306) retinoblastoma binding protein [Rattus norvegicus] >pir [39181 [39181 G1/S transition control protein-binding protein RbAp46 - human >s	(AF103261) immunoglobulin heavy chain variable region [Homo sapiens] Length = 117	HWLRV24R Ig kappa chain Am37 precursor - human Length = 216	HWMCC54R Ig kappa light chain (VJC) [Homo sapiens] >pir S40357 S40357 Ig kappa chain V-J-C region - human {SUB 1-136} >emb CAA61443.1  immunoglobulin anti-F(ab")2 variable region light chain [Homo sapiens] {SUB 21-132} Length = 137	HWLNK85R (AF018265) immunoglobulin lambda light chain [synthetic construct] Length = 236	HCQAV48R IG light chain variable region (VJ) [Homo sapiens] >pir S38643 S38643 lg kappa chain V region - human (fragment) >pir S46369 S46369 lG light chain variable region (VJ) - human {SUB 6-134} >pir A25521 A25521 lg kappa chain V region (321) - human (fragment)	HWMCM79R immunoglobulin kappa-chain [Homo sapiens] Length gb AAA02610.
HCRMF92R	HWLQF89R	HWLOG90R	HWLRV24R	HWMCC54R	HWLNK85R	HCQAV48R	HWMCM79R
2482	2483	2484	2485	2486	2487	2488	2489

HCQAS76 Lambda ZAP	pBluescript	pSport1	pSport1	pSport1	pBluescript	pSport1	pBluescript	pSport1	pSport1	pSport1	pSport1	HCQAM96 Lambda ZAP
HCQAS76	HKLRA71	HWMCJ58	HWLMJ20	HWLMZ25	HKLRB13	HWLMU79	HKLSA25	HWLNN06	HWLMM42	HWMBC38	HWLVU11	нсQам96
76	55	83	79	95	91	95	89	92	75	80	69	50
74	47	20	69	91	85	88	57	74	75	70	69	38
170	223	208	247	423	356	443	477	484	433	455	402	174
3	2	2	2	1	3	3	1	2	242	3	1	-
6767	8929	6929	0229	6771	6772	6773	6774	6775	9229	2229	8229	6179
emb CAA3277 0.1	gb AAB94909. 1	gb AAA02914.	emb CAA6740 6.1	dbj BAA75031 .1	emb CAA0918 3.1	emb CAA6506 1.1	gb AAA71907. 1	gb AAF21612. 1	emb CAA7503 2.1	gb AAC16848. 1	gb AAB00166. 1	gb AAA52938. 1
immunoglobulin variable chain lambda [Homo sapiens] >pir S04519 S04519 Ig lambda chain precursor V-II region (2.1) - human (fragment) Length = 118	mmunoglobulin lambda-2b light chain [Ovis aries] Length = 118	HWMCJ58R IgG [Homo sapiens] Length = 476	immunoglobulin gamma heavy chain [Homo sapiens] Length = 134	loblin heavy chain variable Length = 119	ght chain	immunoglobulin kappa light chain [Homo sapiens] Length = 236	immunoglobulin kappa light chain variable region [Hono sapiens] Length = 106	(AF018265) immunoglobulin lambda light chain [synthetic construct] Length = 236	HWLMM42R immunoglobulin lambda heavy chain [Homo sapiens] >gb AAF14196.1  (AF107231) immunoglobulin heavy chain variable region [Homo sapiens] {SUB 20-147} Length = 477	HWMBC38R (AF063771) immunoglobulin lambda light chain variable region [Homo sapiens] Length = 108	HWLVU11R immunoglobulin lambda-chain subgroup II [Homo sapiens] >dbj BAA20002.1 V1-5 [Homo sapiens] {SUB 20-118} Length = 118	This CDS feature is included to show the translation of the corresponding V_region. Presently translation qualifiers on V_region features are illegal [Homo sapiens] Length = 139
HCQAS76R	HKLRA71R	HWMCJ58R	HWLMJ20R	HWLMZ25R	HKLRB13R	HWLMU79R	HKLSA25R	HWLNN06R	HWLMM42R	HWMBC38R	HWLVUIIR	нсоам96к
2490	2491	2492	2493	2494	2495	2496	2497	2498	2499	2500	2501	2502

HCQCE43 Lambda ZAP	HCQCG90 Lambda ZAP	HCQCK29 Lambda ZAP	HCQCU15 Lambda ZAP	HCQDW78 Lambda ZAP	HCQDW90 Lambda ZAP	pBluescript SK-	pBluescript SK-	HKLAA14 Lambda ZAP	HCQCK49 Lambda ZAP	pSport1	pSport1	pSport1
нсосе43	нсосево	нсоск29	нсоси15	нсорм78	нсорм90	HCYBM34	HCYBM57	HKLAA14	нсоск49	HWLMU27	HWLUR23	HWLRQ41
										97	95	88
										97	95	98
26	2	3	2	24	2	2	9	2	22	139	366	211
133	139	107	76	134	109	109	134	148	114	2	43	53
0829	6781	6782	6783	6784	6785	9829	6787	6788	6429	6790	6791	6792
										gb AAA58453. 1	gb AAA53505. 1	pir A46159 A4 6159
										HWLMU27R initiation factor 4D [Homo sapiens] >gb AAA86989.1  eIF-5A [Homo sapiens] >pir B31486 FHUA translation initiation factor eIF- 5A - human >sp P10159 IF5A_HUMAN INITIATION FACTOR 5A (EIF-5A) (EIF-4D) (REV BINDING FACTOR). {SUB 2-154} Length = 154	insulin-like growth factor binding protein 5 [Homo sapiens] >gb AAA72051.1  [Human insulin-like growth factor binding protein 5 (IGFBP5) gene], gene product [Homo sapiens] >gb AAC09368.1  (AF055033) insulin-like growth factor binding protein 5 [Homo sapie	interferon-dependent positive-acting transcription factor ISGF-3 91K chain - human Length = 739
HCQCE43R	HCQCG90R	HCQCK29R	HCQCU15R	HCQDW78R	HCQDW90R	HCYBM34R	HCYBM57R	HKLAA14R	HCQCK49R	HWLMU27R	HWLUR23R	HWLRQ41R
2503	2504	2505	2506	2507	2508	2509	2510	2511	2512	2513	2514	2515

pSport1	pSport1	HCQDS58 Lambda ZAP	HCQDA89 Lambda ZAP II	HCQCO43 Lambda ZAP II	HCQCG73 Lambda ZAP	pSport1	pSport1	pSport1	pSport1
HWLOC77	HDDNQ21	нсор <b>s</b> 58	нсора89	<b>НС</b> QСО43	керосе 13	НWLQA92	HCROO70	HCRNL15	HCROM41
100	86	99	82	86	26	69	88		86
100	86	28	74	96	62	9	98		86
294	408	384	330	180	328	208	413	233	524
-	-	298	187	-	2	2	252	108	3
6793	6794	6795	9629	6797	8629	6629	0089	6801	6802
gb AAA16521.	emb CAA5933 7.1	gb AAA21551. 1	emb CAA6770 5.1	gb AAA96553. 1	gb AAB48435. 1	dbj BAA13441 .1	dbj BAA04968 .1		dbj BAA06543 .1
HWLOC77R interferon-gamma [Homo sapiens] >gb AAA53230.1  gb AAA16521. interferon-gamma [Homo sapiens]   1    2gb AAF02217.1 AF078829_1 (AF078829) proteasome activator PA28 alpha [Homo sapiens]   2pir A54859 A54859 proteasome activator PA28 alpha chain - human >sp Q06323 IGUP_HUMAN INTE	HDDNQ21R interferon-inducible protein [Homo sapiens] >sp P13164 INI9 HUMAN INTERFERON- INDUCIBLE PROTEIN 9-27 (LEU-13 ANTIGEN). Length = 125	Interleukin 15 [Homo sapiens] >emb CAA62616.1  interleukin-15 [Homo sapiens] >sp P40933 IL15_HUMAN INTERLEUKIN-15 PRECURSOR (IL-15). Length = 162	IP63 protein [Rattus norvegicus] >sp O55160 O55160 IP63 PROTEIN. Length = 571	HCQCO43R J (tail:host specificity;1132) [bacteriophage lambda] Ppir[D43009]QSBPL host specificity protein J - phage lambda >sp[P03749]VHSJ_LAMBD HOST SPECIFICITY PROTEIN J. Length = 1132	KHS1 [Homo sapiens] >splQ9Y4K4 Q9Y4K4 KHS1. Length = 846	HWLQA92R KIAA0007 [Homo sapiens] >sp[Q92577]Q92577 MYELOBLAST KIAA0007 (FRAGMENT). Length = 459	KIAA0036 [Homo sapiens] >sp Q15051 Y036_HUMAN HYPOTHETICAL PROTEIN KIAA0036. Length = 598		KIAA0061 [Homo sapiens] >sp Q15037 Q15037 KIAA0061 PROTEIN (FRAGMENT). Length = 903
HWLOC77R	HDDNQ21R	HCQDS58R	HCQDA89R	нсосо43к	HCQCG73R	HWLQA92R	HCROO70R	HCRNL15R	HCROM41R
2516	2517	2518	2519	2520	2521	2522	2523	2524	2525

pBluescript SK-	pBluescript SK-	HCQDU29 Lambda ZAP	pSport1	pSport1	pBluescript SK-	pSport1	pSport1
H2LAA02	H2CBC43	нсори29	нwмвн25	HWMBW89	HCYBK06	HWMCL19	HCRMR39
86	83		96			86	92
86	82		96			95	65
487	409	73	440	527	148	275	413
122	65	17	8	3	11	KI .	£
6803	6804	6805	9089	6807	8089	6089	6810
dbj BAA09928 .1	gb AAA86132. 1		emb CAA3203			emb CAA5112 0.1	gb AAA85268. 1
KIAA0158 gene product is related to Drosophila DiffG protein. [Homo sapiens] >gb AAB92377.1  (AF038404) homolog of Nedd5; hNedd5 [Homo sapiens] >sp Q15019 NED5_HUMAN NEDD5 PROTEIN HOMOLOG (KIAA0158). >gb AAD12225.1  (AC005104) KIAA0158; similar to human N	kinesin-like spindle protein HKSP [Homo sapiens]   2pir G02157 G02157 kinesin-like spindle protein HKSP - human Length = 1056		HWMBH25R lactate dehydrogenase B [Homo sapiens] emb   Semb CAA68701.1  lactate dehydrogenase B (AA 1 - 3.1  334) [Homo sapiens] > pir S02795 DEHULH L-lactate dehydrogenase (EC 1.1.1.27) chain H - human   Sep P07195 LDHH HUMAN L-LACTATE   DEHYDROGENASE H CHAIN (EC 1.1.1.27) (LDH-B			HWMCL19R Ig kappa light chain (VJC) [Homo sapiens]  >emb[CAB75876.1] (AJ272080) immunoglobulin light chain variable region [Homo sapiens] {SUB 14-131} >gb[AAD16547.1] (AF103376) immunoglobulin kappa light chain variable region [Homo sapiens] {SUB 22-122} >pir[S3409	lumican [Homo sapiens] >sp P51884 LUM_HUMAN  gb AAA85268. LUMICAN PRECURSOR (LUM) (KERATAN 1  SULFATE PROTEOGLYCAN). Length = 338
H2LAA02R	H2CBC43R	нсори29к	HWMBH25R	HWMBW89R	HCYBK06R	HWMCL19R	HCRMR39R
2526	2527	2528	2529	2530	2531	2532	2533

pSport1	pSport1	pSport1	pSport1	pSport1
HWMBJ73	HWLVE15	HCRMD64	HCRNO44	HNBTK71
88	96	96	78	16
88	96	96	78	91
428	392	185	514	628
3	E.	r,	299	2
6811	6812	6813	6814	6815
gb AAA59599. 1	emb CAA4211 8.1	gb AAA91780. 1	emb CAB5593	gb AAA59982.
HWMBJ73R [lymphocyte antigen [Homo sapiens] >gb AAA59613.1  HLA-A31 precursor [Homo sapiens] >gb AAB05976.1  Jymphocyte antigen [Homo sapiens] >pir I72170 I72170 MHC class I histocompatibility antigen HLA-A31 alpha chain (allele A*31012) precursor - human >sp P1618	HWLVE15R M1 subunit of ribonucleotide reductase [Homo sapiens] >emb CAA42180.1  large subunit ribonucleotide reductase [Homo sapiens] >pir S16680 S16680 ribonucleoside-diphosphate reductase (EC 1.17.4.1) chain M1 - human >sp P23921 RIR1_HUMAN RIBONUCLEOSIDE-DIPHOS	_ Z	HCRNO44R (AL117452) hypothetical protein [Homo sapiens] >emb[CAB55934.1] (AL117452) hypothetical protein [Homo sapiens] >pir[T17244[T17244 hypothetical protein DKFZp586G1517.1 - human (fragment) >sp[CAB55934 CAB55934 Hypothetical 99.4 kd protein (fragment). >emb[C	membrane protein [Homo sapiens]  >emb[CAA42708.1  MRP-1 (motility related protein) 1  [Homo sapiens] >gb AAA80320.1  CD9 antigen [Homo sapiens] >gb AAC60586.1  CD9 antigen [Ihuman, leukocytes, Peptide, 228 aa] [Homo sapiens]  >pir A46123 A40402 CD9 antigen - hu
HWMBJ73R	HWLVE15R	HCRMD64RA	HCRN044R	HNBTK71R
2534	2535	2536	2537	2538

pSport1	pSport1	pSport1	HCQDI65 Lambda ZAP II	pSport1	pSportl
HCROL22	HWLVI52	HSAMD89	нсдрі65	HCROE42	HCROJ80
47	100	75	96	82	89
45	100	73	96	08	89
371	413	476	222	551	475
٤	<i>c</i>	6	112	£	149
6816	6817	6818	6819	6820	6821
dbj BAA11528	gb AAA59581. 1	gb AAA21805.	gb AAA78807.	gb AAD56542. 1 AF1843	gb AAC51231. 1
membrane protein with histidine rich charge clusters dbj BAA11528 [Homo sapiens] >gb AAD12305.1  (AF117221) KE41  protein [Homo sapiens] >sp Q92504 Q92504 MEMBRANE PROTEIN WITH HISTIDINE RICH CHARGE CLUSTERS. Length = 429	metalloproteinase inhibitor precursor [Homo sapiens] gb AAA59581.  >gb AAA61186.1  metalloproteinase-2 inhibitor precursor [Homo sapiens] >gb AAB19474.1  tissue inhibitor of metalloproteinase 2, TIMP-2 {EC 3.4.24} [human, Peptide, 220 aa] [Homo sapiens] >pir A37128 A	non-muscle alpha tropomyosin [Rattus norvegicus] >sp Q63582 Q63582 NON-MUSCLE ALPHA TROPOMYOSIN. Length = 284	mitochondrial ATP synthase subunit 9 precursor [Homo sapiens] >pir 138612 138612 H+-transporting ATP synthase (EC 3.6.1.34) lipid-binding protein P3 precursor, mitochondrial - human >sp P48201 AT93_HUMAN ATP SYNTHASE LIPID-BINDING PROTEIN P3 PRECURSOR (EC	(AF184344) DNA polymerase accessory subunit precursor [Homo sapiens] >sp AAD56542 AAD56542 DNA polymerase accessory subunit precursor. Length = 485	mitochondrial intermediate peptidase precursor [Homo sapiens] >sp[Q99797]PMIP_HUMAN MITOCHONDRIAL INTERMEDIATE PEPTIDASE PRECURSOR (EC 3.4.24.59) (MIP). Length = 713
HCROL22R	HWLVI52R	HSAMD89R	HCQDI65R	HCROE42R	HCROJ80R
2539	2540	2541	2542	2543	2544

ZAP Express	pSport1	pSport1	pSport1	Lambda ZAP II	pSport1	pSport1
HCWHT65 ZAP Express	HCRMX32	HCROE77	HCRMX69	нсорн45	HOCTA19	HWLRA67
74		93	93	88	92	78
70		93	93	98	80	89
432	300	329	144	312	504	117
	1	£	-	106	217	-
6822	6823	6824	6825	6826	6827	6828
gb AAC51231. 1		gb AAA67526. 1	dbj BAA11423	emb CAA4767	gb AAA65087.	emb CAA3820 1.1
HCWHT65R mitochondrial intermediate peptidase precursor [Homo sapiens] >sp Q99797 PMIP_HUMAN MITOCHONDRIAL INTERMEDIATE PEPTIDASE PRECURSOR (EC 3.4.24.59) (MIP). Length = 713		MTHSP75 [Homo sapiens] >sp P38646 GR75_HUMAN MITOCHONDRIAL STRESS-70 PROTEIN PRECURSOR (75 KD GLUCOSE REGULATED PROTEIN) (GRP 75) (PEPTIDE-BINDING PROTEIN 74) (PBP74) (MORTALIN) (MOT). Length = 679	HCRMX69R multifunctional protein CAD [Homo sapiens] >sp P27708 PYR1 HUMAN CAD PROTEIN [INCLUDES: GLÜTAMINE-DEPENDENT CARBAMOYL-PHOSPHATE SYNTHASE (EC 6.3.5.5), ASPARTATE CARBAMOYLTRANSFERASE (EC 2.1.3.2); DIHYDROOROTASE (EC 3.5.2.3)]. >gb AAA51907.1  CAD [Homo sap	myosin I heavy chain isoform [Gallus gallus] >spjQ02440 MYSD_CHICK DILUTE MYOSIN HEAVY CHAIN, ISOFORM I (MYOSIN HEAVY CHAIN P190) (MYOSIN-V). Length = 1829	myosin II nonmuscle [Rana catesbeiana] >sp Q91304 Q91304 MYOSIN II NONMUSCLE (FRAGMENT). Length = 261	myosin regulatory light chain [Homo sapiens] >pir S11493 MOHULP myosin regulatory light chain, placental - human >sp P19105 MLRM_HUMAN MYOSIN REGULATORY LIGHT CHAIN 2, NONSARCOMERIC (MYOSIN RLC). {SUB 2-171} Length = 171
HCWHT65R	HCRMX32R	HCROE77R	HCRMX69R	<b>НСQDH</b> 45R	HOCTA19R	HWLRA67R
2545	2546	2547	2548	2549	2550	2551

pSport1	pBluescript SK-	pSport1	HFPBS29 Uni-ZAP XR	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II
HWLOM88	H2CBI14	HCRNI08	HFPBS29	нсорг20	НСQС <b>В</b> 43	<b>НС</b> QDA51
70	92	88	100		43	43
69	09	88	97		40	41
409	208	537	183	226	113	322
2	2	-		128	27	260
6859	6830	6831	6832	6833	6834	6835
gb AAF36524. 1 AF1320	emb CAA6903 6.1	dbj BAA04570	emb CAA5222		dbj BAA07291 .1	dbj BAA07291 .1
HWLOM88R (AF132021) myosin X [Homo sapiens] >gb AAF17363.1 AF184153_1 (AF184153) myosin X [Homo sapiens] {SUB 347-495} Length = 1540	mysoin heavy chain 12 [Homo sapiens] >sp CAA69036 CAA69036 Mysoin heavy chain 12. Length = 1828		NAD+-isocitrate dehydrogenase, gamma subunit [Macaca fascicularis] >pir S39065 S39065 isocitrate dehydrogenase (NAD+) (EC 1.1.1.41) gamma chain precursor - crab-eating macaque (fragment) >sp P41564 IDHG_MACFA ISOCITRATE DEHYDROGENASE [NAD] SUBUNIT GAMMA,		HCQCB43R NADH dehydrogenase subunit 2 [Homo sapiens] >dbj BAA07291.1  NADH dehydrogenase subunit 2 [Homo sapiens] >sp Q34769 Q34769 NADH DEHYDROGENASE SUBUNIT 2. Length = 347	NADH dehydrogenase subunit 2 [Homo sapiens] >dbj BAA07291.1  NADH dehydrogenase subunit 2 [Homo sapiens] >sp Q34769 Q34769 NADH DEHYDROGENASE SUBUNIT 2. Length = 347
HWLOM88R	H2CBI14R	HCRNI08R	HFPBS29R	HCQDL50R	HCQCB43R	HCQDA51R
2552	2553	2554	2555	2556	2557	2558

Lambda ZAP II	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II	HCQCT16 Lambda ZAP	HCQDA65 Lambda ZAP
HCQDB27 Lambda ZAP	HCQDS85	HCQCR82	HCQDV94   Lambda ZAP	нсост16	нсорде <b>6</b> 5
63	49	100	75		68
28	43	06	72		68
109	451	206	323	49	457
23	284	84	174	2	2
6836	6837	6838	6839	6840	6841
dbj BAA07291	dbj BAA07291 .1	.1	dbj BAA77673 .1		emb CAA5812 7.1
HCQDB27R NADH dehydrogenase subunit 2 [Homo sapiens] >dbj BAA07291.1  NADH dehydrogenase subunit 2 [Homo sapiens] >sp Q34769 Q34769 NADH DEHYDROGENASE SUBUNIT 2. Length = 347	HCQDS85R NADH dehydrogenase subunit 2 [Homo sapiens] >dbj BAA07291.1  NADH dehydrogenase subunit 2 [Homo sapiens] >sp Q34769 Q34769 NADH DEHYDROGENASE SUBUNIT 2. Length = 347	NADH dehydrogenase subunit 3 [Homo sapiens] >dbj BAA77672.1  NADH dehydrogenase subunit 3 [Homo sapiens] >sp BAA77672 BAA77672 NADH dehydrogenase subunit 3. >gb AAB63453.1  (AF004342) NADH dehydrogenase III [Homo sapiens] {SUB 10-115} >dbj BAA76519.1  (AB	HCQDV94R NADH dehydrogenase subunit 4 [Homo sapiens] >dbj BAA77673.1 NADH dehydrogenase subunit 4 [Homo sapiens] >sp BAA77673 BAA77673 NADH dehydrogenase subunit 4. Length = 459		neutrophil gelatinase associated lipocalin [Homo sapiens] >sp P80188 NGAL_HUMAN NEUTROPHIL GELATINASE-ASSOCIATED LIPOCALIN PRECURSOR (NGAL) (P25) (25 KD ALPHA-2-MICROGLOBULIN-RELATED SUBUNIT OF MMP-9) (LIPOCALIN-2) (ONCOGENE 24P3). >gb AAD14168.1 575256_1
нсдрв27R	HCQDS85R	HCQCR82R	<b>НСQDV94R</b>	HCQCT16R	нсорабя
2559	2560	2561	2562	2563	2564

pSport1	pBluescript SK-	pSport1	pBluescript SK-	pSport1	pSport1	pBluescript SK-
<b>Н</b> WLWH33	HCYBJ83	HWLRE17	H2LAC53	HWLOM10	HWLQ029	H2LBA48
91	98	73	87	76	9/	63
06	81	73	98	96	9/	41
547	494	415	399	438	332	432
200	285	2	-	-	3	172
6842	6843	6844	6845	6846	6847	6848
emb CAA5812 7.1	gb AAB99856.	gb AAA35537.	gb AAA36380. 1	gb AAA59954. 1	dbj BAA01980	emb CAA9590 7.1
HWLWH33R neutrophil gelatinase associated lipocalin [Homo sapiens] >sp P80188 NGAL_HUMAN NEUTROPHIL GELATINASE-ASSOCIATED LIPOCALIN PRECURSOR (NGAL) (P25) (25 KD ALPHA-2-MICROGLOBULIN-RELATED SUBUNIT OF MMP-9) (LIPOCALIN-2) (ONCOGENE 24P3). >gb AAD14168.1 S75256_1	(AF043542) nucleoside diphosphate kinase [Gallus gallus] >sp O57535 O57535 NUCLEOSIDE DIPHOSPHATE KINASE. Length = 153	nuclear autoantigen [Homo sapiens] >pir A37244 A37244 nuclear autoantigen Sp-100 - human Length = 480	nucleophosmin [Homo sapiens] >gb AAA36385.1  nucleolar protein B23 [Homo sapiens] >gb AAA58386.1  nucleolar phosphoprotein B23 [Homo sapiens] >gb AAB94739.1  nucleophosmin phosphoprotein [Homo sapiens] >pir A33423 A32915 nucleophosmin - human	nucleolin [Homo sapiens] >pir A35804 A35804 nucleolin - human >sp P19338 NUCL_HUMAN NUCLEOLIN (PROTEIN C23). {SUB 2-707} Length = 707	ORF [Pan troglodytes] >sp Q28808 INI2_PANTR INTERFERON-INDUCED PROTEIN 6-16 PRECURSOR (IFI-6-16). Length = 130	ORF YNL040w [Saccharomyces cerevisiae] >pir S62962 S62962 hypothetical protein YNL040w - yeast (Saccharomyces cerevisiae) >sp P53960 YNE0_YEAST HYPOTHETICAL 51.0 KD PROTEIN IN YIP3-TFC5 INTERGENIC REGION. Length = 456
HWLWH33R	HCYBJ83R	HWLRE17R	H2LAC53R	HWLOM10R	HWLQ029R	H2LBA48R
2565	2566	2567	2568	2569	2570	2571

4 659 38 64 HCRPZ16 pSport1	5 3 66 69 HKCSA80 pBluescript	145 54 72 HCQBD02 Lambda ZAP	193 97 97 HCRPH64 pSport1	319 83 83 HDTBZ03 pCMVSport 2.0	6 347 HLYED39 pSport1	246 97 98 HCQCB85 Lambda ZAP	130 53 56 HCRME24 pSport1	5 239 63 73 HWLQK64 pSport1
6849 294	6850 245	6851 2	6852 2	6853 2	6854 216	6855 1	6856 35	6857 445
	gb AAA72122. 68	emb CAA3648 68 0.1		gb AAC39523. 68 1	9	9.1	gb AAA60065. 68	
HCRPZ16R (AL033502) hypothetical protein [Candida albicans] emb CAA2200 >sp O94058 O94058 HYPOTHETICAL 85.6 KD   9.1  PROTEIN. Length = 747	ORF1 [Escherichia coli] Length = 334	ORFII [Homo sapiens] >splQ14754 Q14754 ORFII. Length = 712	ornithine decarboxylase antizyme [Homo sapiens] Length = 228	HDTBZ03R OS9 [Homo sapiens] Length = 474		peptidylprolyl isomerase [Homo sapiens] >emb CAA68264.1  cyclophilin (AA 1-165) [Homo sapiens] >gb AAB81959.1  (AF023859) cyclophilin A [Papio hamadryas] >gb AAB81960.1  (AF023860) cyclophilin A [Cercopithecus aethiops] >gb AAB81961.1  (AF023861) cyclophi	HCRME24R perforin [Homo sapiens] >gb AAA60167.1  perforin [Homo sapiens] >pir A45816 A37181 perforin 1 precursor - human >sp P14222 PERF_HUMAN PERFORIN 1 PRECURSOR (P1) (LYMPHOCYTE PORE FORMING PROTEIN) (PFP) (CYTOLYSIN). Length = 555	HWLQK64R PEX5p [Mus musculus] >sp 009012 PEX5_MOUSE emb CAB0969 PEROXISOMAL TARGETING SIGNAL 1 RECEPTOR (PEROXISMORE RECEPTOR 1) (PEROXISOMAL C-TERMINAL TARGETING SIGNAL IMPORT RECEPTOR) (PTS1-BP) (PEROXIN-5) (PTS1 RECEPTOR) (PXR1P)
HCRPZ16R	HKCSA80R	HCQBD02R	HCRPH64R	HDTBZ03R	HLYED39R	нсосвяя	HCRME24R	HWLQK64R
2572	2573	2574	2575	2576	2577	2578	2579	2580

pSport1	HCQDL14 Lambda ZAP	pSport1	pBluescript SK-	Lambda ZAP II	pSport1	pSport1	pSport1	pBluescript SK-
HCRNF48	нсорг14 1	HWLQA11	нсув <i>н</i> 73	HCQCJ88	HWLXJ34	HOCTB09	HCRQN67	нсувн30
78	83	72	78	88	98	95	100	82
78	77	69	75	88	83	92	100	79
383	339	257	269	272	416	477	260	450
m	-	48	6	٣	3	-	ဂ	292
8858	6859	0989	6861	6862	6863	6864	6865	6866
emb CAA4264	gb AAA31441. 1	pir A61382 A6 1382	emb CAA6626 5.1	gb AAA59972.	gb AAC00205. 1	gb AAA51889. 1	emb CAA6826	gb AAF14864. 1 AF1131
HCRNF48R phosphate carrier protein [Homo sapiens] >emb CAB56612.1  phosphate carrier [Homo sapiens] >pir B53737 B53737 phosphate carrier protein, form B - human >sp CAB56612 CAB56612 Phosphate carrier. Length = 361	HCQDL14R phosphofructokinase [Oryctolagus cuniculus]  >pir[A26550]KIRBF 6-phosphofructokinase (EC 2.7.1.11), muscle - rabbit >sp]P00511]K6PF_RABIT 6-PHOSPHOFRUCTOKINASE, MUSCLE TYPE (EC 2.7.1.11) (PHOSPHOFRUCTOKINASE 1) (PHOSPHOHEXOKINASE) (PHOSPHOFRUCTO- 1-KINASE I	phosphorylation regulatory protein HP-10 - human Length = 492	plakophilin 2a [Homo sapiens] >sp Q99960 Q99960 PLAKOPHILIN 2A. Length = 837	fusion protein [Homo sapiens] >gb AAD13865.1 1680464_1 promyelocytic leukemia protein [Homo sapiens] {SUB 220-333} Length = 744	PRAJA1 [Mus musculus] >sp 055176 055176 PRAJA1. Length = 424	C9 complement protein [Homo sapiens] Length = 557	precursor polypeptide (AA -23 to 1120) [Homo sapiens] >gb AAD15273.1  T200 glycoprotein [Homo sapiens] >sp Q16614 Q16614 T200 LEUKOCYTE COMMON ANTIGEN (CD45, LC-A) PRECURSOR (EC 3.1.3.48) (CD45, LC-A). Length = 1143	(AF113123) carbonyl reductase [Homo sapiens] >sp AAF14864 AAF14864 Carbonyl reductase. Length = 244
HCRNF48R	HCQDL14R	HWLQA11R	нсувн738	HCQC188R	HWLXJ34R	HOCTB09R	HCRQN67R	HCYBH30R
2581	2582	2583	2584	2585	2586	2587	2588	2589

			Ħ		
pSport1	pSport1	pSport1	pCMVSport 2.0	pSport1	pSport1
HCRN004	HWLRC47	HCROE26	HOHBE57	HCROK02	HWMBB94
96		98	84	86	100
96		98	9/	86	86
104	236	260	535	310	272
3	3	8	59	2	81
6867	8989	6989	0890	6871	6872
gb AAA52129.		gb AAB32370.	emb CAA7048 8.1	emb CAA3678	gb AAA60222. 1
HCRNO04R preprocathepsin B [Homo sapiens] >pir A26498 KHHUB cathepsin B (EC 3.4.22.1) precursor - human >sp P07858 CATB_HUMAN CATHEPSIN B PRECURSOR (EC 3.4.22.1) (CATHEPSIN B1) (APP SECRETASE). Length = 339		prostacyclin-stimulating factor, PGI2-stimulating factor, PSF [human, cultured diploid fibroblast cells, Peptide, 282 aa] [Homo sapiens] >pir S50031 S50031 prostacyclin-stimulating factor - human >sp Q16270 Q16270 PROSTACYCLIN-STIMULATING FACTOR. Length =	protein kinase [Homo sapiens] >sp[Q92631 Q92631 PROTEIN KINASE (FRAGMENT). Length = 240	protein phophatase 2A alpha catalytic subunit (AA 1- emb CAA3678 309) [Bos taurus] >emb CAA51381.1  protein phosphatase-2A [Bos taurus] >gb AAA30981.1  protein phosphatase 2A alpha subunit [Sus scrofa] >emb CAA29471.1  phosphatase (AA 1-309) [Oryctolagus cuniculus] >g	protein tyrosine phosphatase [Homo sapiens] >gb AAA66496.1  protein phosphatase [Homo sapiens] >sp Q16667 CDN3_HUMAN CYCLIN-DEPENDENT KINASE INHIBITOR 3 (EC 3.1.3.48) (EC 3.1.3.16) (CDK2- ASSOCIATED DUAL SPECIFICITY PHOSPHATASE) (KINASE ASSOCIATED PHOSPHA
HCRNO04R	HWLRC47R	HCROE26R	нонве57R	HCROK02R	HWMBB94R
2590	2591	2592	2593	2594	2595

HUVHA17 Uni-ZAP XR	Uni-ZAP XR	pSport1	pBluescript	Lambda ZAP II	pSportl	Uni-ZAP XR	HOSBE19 Uni-ZAP XR	pSport1
HUVHA17	HLTI191	HCRMC40	HKCSL44	HCQAR83	нwLQD31	HOUDN78 Uni-ZAP XR	HOSBE19	HCROB08
95		100	69	95	97	52	74	
91		100	99	93	97	84	5	
443	230	160	288	184	250	307	367	170
174	3	2	37	11	2	2	161	42
6873	6874	6875	9289	6877	8289	6289	0889	6881
gb AAA37592.  1		emb CAA3097 6.1	gb AAA36021. 1	gb AAB41848. 2	gb AAA36563.	emb CAA8162 6.1	dbj BAA11211 .1	
HUVHA17R focal adhesion kinase [Mus musculus] >pir A46166 A46166 protein-tyrosine kinase (EC 2.7.1.112) - mouse >sp P34152 FAK1_MOUSE FOCAL ADHESION KINASE 1 (EC 2.7.1.112) (FADK 1) (PP125FAK). Length = 1052		put. ORF [Homo sapiens] >pir A31026 A31026 probable membrane receptor protein - human >sp P08910 HPS1_HUMAN PROTEIN PHPS1-2. Length = 425	Q1Z 7F5 [Homo sapiens] >gb AAA36378.1  may code for Wilm"s tumor-related protein [Homo sapiens] >gb AAA63253.1  Wilm"s tumor-related protein [Homo sapiens] >gb AAB27665.1  QM [human, nontumorigenic Wilms" microcell hybrid cells, Peptide, 214 aa] [Homo	sperm membrane protein BS-63 [Homo sapiens] gb AAB41848. >sp AAB41848 AAB41848 Sperm membrane protein 2  BS-63. >sp Q99666 Q99666 BS-63. {SUB 1197-1765} Length = 1765	rapamycin- and FK506-binding protein [Homo sapiens] >pir JC1365 JC1365 FK506/rapamycin-binding protein FKBP13 precursor - human Length = 142	HOUDN78R Rab5c protein [Canis familiaris] >pir S65933 S65933 emb CAA8162 GTP-binding protein Rab5c - dog   5xp P51147 RB5C CANFA RAS-RELATED   PROTEIN RAB-5C. Length = 216	ras-related GTP-binding protein [Homo sapiens] Length = 184	
HUVHA17R	HLTIJ91R	HCRMC40R	HKCSL44R	HCQAR83R	HWLQD31R	HOUDN78R	HOSBE19R	HCROB08R
2596	2597	2598	2599	2600	2601	2602	2603	2604

pSport1	pSport1	pSport1	pSport1	pSport
HWLQG37	HSAMB82	HWLWE05	HWLRB68	HWLMB86
HMI	HSA	HWI	HM	HWI
09	68		100	88
09	68	92	100	88
387	311	323	465	149
205	23	18	_	39
6882	6883	6884	6885	9889
emb CAA8148	gb AAC15856.	emb CAA6158	gb AAA03341.	gb AAA19815.
8.1	gb AA(	emb C/ 2.1	gb AA.	gb AA/
HWLQG37R ribosomal protein [Homo sapiens]  >emb CAA40328.1  ribosomal protein L38 [Rattus rattus] >pir S15658 R5RT38 ribosomal protein L38 - rat >pir S38385 S38385 ribosomal protein L38 - human >sp P23411 RL38 HUMAN 60S RIBOSOMAL PROTEIN L38. {SUB 2-70} >dbj BAA258	ribosomal protein L11 [Homo sapiens]  >emb CAA44072.1  ribosomal protein L11 [Rattus 1 rattus] >pir S17351 R5RT11 ribosomal protein L11 precursor - rat >sp P39026 RL11 HUMAN 60S RIBOSOMAL PROTEIN L11. {SUB 2-178} >dbj BAA25831.1  (AB007171) ribosomal protein	ribosomal protein L21 [Homo sapiens]  >gb AAA85655.1  ribosomal protein L21 [Homo 2apiens] >pir S55913 S55913 ribosomal protein L21, cytosolic - human >sp P46778 RL21 HUMAN 60S RIBOSOMAL PROTEIN L21. {SUB 2-160}  >dbj BAA25835.1  (AB007176) ribosomal protei	ribosomal protein L23a [Homo sapiens] >gb AAA35681.1  homology to rat ribosomal protein 1 L23 [Homo sapiens] {SUB 10-156} Length = 156	ribosomal protein L27 [Homo sapiens] >gb AAC15857.1  ribosomal protein L27 [Homo sapiens] >emb CAA30313.1  ribosomal protein L27 (AA 1 - 136) [Rattus norvegicus] >gb AAF25951.1 AF214527_1 (AF214527) ribosomal protein L27 [Mus musculus] >emb CAA40181.1  ri
HWLQG37R	HSAMB82R	HWLWE05R	HWLRB68R	HWLMB86R
2605	2606	2607	2608	5609

pSport1	HAIDT43 Uni-ZAP XR	pBluescript SK-	НСQDU05   Lambda ZAP II	pSport1	pSport1
100 HWLQB60	HAIDT43	H2LAU86	нсорооз	HCRPM16	нwLQA31
100		66	96	100	06
100		86	96	100	87
430	140	477	197	362	240
2	m		m	<del>س</del>	-
6887	8889	6889	0689	6891	6892
emb CAA5284  8.1		gb AAC41916. 1	gb AAB65437. 1	emb CAA4934 5.1	gb AAB06757. 1
HWLQB60R ribosomal protein L28 [Mus musculus] >pir[148738 148738 ribosomal protein L28 - mouse >sp P41105 RL28_MOUSE 60S RIBOSOMAL PROTEIN L28. {SUB 2-137} Length = 137		ribosomal protein L34 [Homo sapiens] >pir 168524 168524 ribosomal protein L34 - human >sp P49207 RL34_HUMAN 60S RIBOSOMAL PROTEIN L34. {SUB 2-117} Length = 117	HCQDU05R (AF013215) ribosomal protein S2 [Bos taurus] >sp O18789 RS2_BOVIN 40S RIBOSOMAL PROTEIN S2 (FRAGMENT). Length = 286	HCRPM16R ribosomal protein S26 [Homo sapiens]  >dbj BAA25824.1  (AB007161) ribosomal protein S26 [Homo sapiens] {SUB 62-106}  >emb CAA55818.1  ribosomal protein S26 [Homo sapiens] {SUB 1-20} >dbj BAA25823.1   (AB007160) ribosomal protein S26 [Homo sapiens] {SUB 38-60	HWLQA31R ribosomal protein S29 [Bos taurus] >gb AAA85661.1  ribosomal protein S29 [Homo sapiens] >gb AAB27426.1  homologous to antisense sequence of krev-1, anti oncogene [Homo sapiens] >emb CAA41778.1  ribosomal protein S29 [Rattus norvegicus] >gb AAB27429.1  S29
HWLQB60R	HAIDT43R	H2LAU86R	нсороозк	HCRPM16R	HWLQA31R
2610	2611	2612	2613	2614	2615

pBluescript	pBluescript	pBluescript	pSport1	pSport1	Lambda ZAP II	pSport1
HFVKA92	HKLSA82	HKLSA88	HWLNK27	HCRNT24	HCQAW95	HWLMP89
87	70	100	85	81	94	84
83	29	100	84	81	85	82
257	296	246	345	299	474	328
<del>د</del>	3	127	34	27	_	2
6893	6894	6895	9689	2689	8689	6899
gb AAA60289.	gb AAC41754. 1	gb AAB08488. 2	gb AAA59512. 1	emb CAA4656 6.1	gb AAA95995. 1	gb AAA65596. 1
HFVKA92R ribosomal protein S6 [Homo sapiens] >emb CAA47719.1  ribosomal protein S6 [Homo sapiens] >gb AAA42079.1  ribosomal protein S6 [Rattus norvegicus] >emb CAA68430.1  ribosomal protein S6 [Mus musculus] >emb CAA90936.1  rpS6 [Mus musculus] >pirJJC1394 R3HU6 r	Rieske Fe-S protein [Homo sapiens] >sp P47985 UCRI_HUMAN UBIQUINOL- CYTOCHROME C REDUCTASE IRON-SULFUR SUBUNIT, MITOCHONDRIAL PRECURSOR (EC 1.10.2.2) (RIESKE IRON-SULFUR PROTEIN) (RISP). Length = 274	replication protein A complex 34 kd subunit homolog Rpa4 [Homo sapiens] >sp AAB08488[AAB08488 Replication protein A complex 34 kd subunit homolog Rpa4. Length = 261	S-lac lectin [Homo sapiens] >gb AAA59513.1  S-lac lectin [Homo sapiens] >emb CAB42834.1  (AL022315) dJ117715.3 (Lectin, Galactose-binding, soluble, 2 (Galectin 2, S-Lac Lectin 2, HL14)) [Homo sapiens] >pir A38140 A38140 galectin 2 - human >sp P05162 LEG2_	S100P calcium-binding protein [Homo sapiens] >pir[S24146[S24146 S-100 protein P - human >sp[P25815[S10E_HUMAN S-100P PROTEIN. Length = 95	similar to product encoded by GenBank Accession number S62516 [Rattus norvegicus] >sp Q62742[Q62742 SA (FRAGMENT). Length = 106	sarcolemmal associated protein-2 [Oryctolagus cuniculus] >sp Q28622 Q28622 SARCOLEMMAL ASSOCIATED PROTEIN-2. Length = 402
HFVKA92R	HKLSA82R	HKLSA88R	HWLNK27R	HCRNT24R	HCQAW95R	HWLMP89R
2616	2617	2618	2619	2620	2621	2622

pSport1	pBluescript	Uni-ZAP XR	HCQCQ84 Lambda ZAP	pSport1	pSport1
HWLVA90	HKCSI14	HFCES53	нсосо <sub>84</sub>	HWLMV10	HWMBC92
94	96		94	78	96
98	96		94	78	96
278	95	165	404	155	268
<u>س</u>	E.	-	8	m	2
0069	6901	6902	6903	6904	6905
emb CAA2521	gb AAA59981. 1		gb AAC51784. 1	emb CAA4700	gb AAB94646. 1
HWLVA90R SB beta-chain (1 is 2nd base in codon) [Homo sapiens] >sp Q14465 Q14465 SB BETA-CHAIN (CLONE PII-BETA-7) (FRAGMENT). >gb AAA36311.1  MHC HLA-SB beta chain [Homo sapiens] {SUB 48-234} >gb AAA59746.1  MHC DP-beta, allele DPB7 [Homo sapiens] {SUB 1-87} Lengt	secretory protein [Homo sapiens] >gb AAA83628.1  intestinal trefoil factor [Homo sapiens] >pir A48284 A48284 intestinal trefoil factor 3 precursor - human >sp Q07654 ITF_HUMAN INTESTINAL TREFOIL FACTOR PRECURSOR (HP1.B). Length = 80		serine protease [Homo sapiens] >sp O15393 TMS2_HUMAN TRANSMEMBRANE PROTEASE, SERINE 2 (EC 3.4.21). Length = 492	HWLMV10R serine/threonine protein kinase [Homo sapiens] em >pir S32831 S32831 serine/threonine-specific protein 5.1 kinase PCTAIRE-3 (EC 2.7.1) - human (fragment) >sp Q07002 KPT3_HUMAN SERINE/THREONINE PROTEIN KINASE PCTAIRE-3 (EC 2.7.1) (FRAGMENT). Length = 380	HWMBC92R integrin binding protein kinase [Mus musculus] >sp[O55222 O55222 INTEGRIN LINKED KINASE (INTEGRIN BINDING PROTEIN KINASE). Length = 452
HWLVA90R	HKCSI14R	HFCES53R	HCQCQ84R	HWLMV10R	HWMBC92R
2623	2624	2625	2626	2627	2628

HBJMM52 Uni-ZAP XR	pSport1	pSport1	pSport1	pCMVSport 3.0	pSport1	HCQDT79 Lambda ZAP	Lambda ZAP II	pSport1
HBJMM52	HCROZ52	НWLQQ35	HHMMF20	ннеиw25	HCRNZ02	нсорт79	нсорг92	HWLVG33
74		92		86	91	92		89
73		75		86	68	84		<i>L</i> 9
333	115	413	101	336	344	361	152	377
-	2	3	3	T	es .	119	3	m
9069	2069	8069	6069	0169	6911	6912	6913	6914
emb CAA6263 5.1		dbj BAA11481 .1		gb AAB58251. 1	gb AAB48981. 1	dbj BAA74949 .1		gb AAA16315. 1
HBJMM52R seryl-tRNA synthetase [Homo sapiens] >pir G01026 G01026 serinetRNA ligase (EC 6.1.1.11) - human >sp P49591 SYS_HUMAN SERYL-TRNA SYNTHETASE (EC 6.1.1.11) (SERINETRNA LIGASE) (SERRS). >pir S00490 S00490 RNA-binding protein, 62K - rabbit (fragment) {SUB		similar to human DNA-binding protein 5. [Homo db sapiens] >sp Q14673 Q14673 KIAA0164 PROTEIN1  Length = 920		similar to mouse Int-6 [Homo sapiens]  >gb AAB88873.1  Int-6 [Homo sapiens]  >gb AAC51760.1  eIF3-p48 [Homo sapiens]  >gb AAC51919.1  mammary tumor-associated protein INT6 [Homo sapiens]  >sp Q64252 IF36_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNI	sodium channel 2 [Homo sapiens]   gb AAB48981.  >sp P78349 P78349 SODIUM CHANNEL 2. Length   1  = 528	Son of sevenless 1 [Rattus norvegicus] >sp[Q9Z111 Q9Z111 SON OF SEVENLESS 1 (FRAGMENT). Length = 204		splicing factor [Homo sapiens] >emb CAA53512.1  gClq-R [Homo sapiens] >pir JT0762 JT0762 pre- mRNA splicing factor SF2 P32 chain precursor - human >sp Q07021 MA32_HUMAN COMPLEMENT COMPONENT 1, Q SUBCOMPONENT BINDING PROTEIN, MITOCHONDRIAL PRECURSOR (GLYCOP
HBJMM52R	HCROZ52R	HWLQQ35R	HHMMF20R	HHEUW25R	HCRNZ02R	нсорт798	HCQDL92R	HWLVG33R
2629	2630	2631	2632	2633	2634	2635	2636	2637

pSport1	HCQDW65 Lambda ZAP	HCQCV70 Lambda ZAP	HCQDN27 Lambda ZAP	Lambda ZAP II	pSport1	pSport1	pSport1	pBluescript SK-	pBluescript SK-
HCR0013	нсорм65	нсосу70	HCQDN27	нсос192	HCRMP36	HCROV67	HCROT79	H2CAA07	H2LAD20
72	93	91	93	91	86	62	100	25	29
51	92	81	91	06	95	55	100	35	57
373	408	162	448	220	353	397	091	360	258
2	40	-	143	2	3	5	2	-	130
6915	6916	6917	6918	6169	6920	6921	6922	6923	6924
gb AAF18954. 1 AF1645	dbj BAA91818 .1	dbj BAA33580	gb AAA84389. 1	emb CAA5183 9.1	emb CAA3962 8.1	gb AAA49087.  1	gb AAD24668. 1 U86074	emb CAA1002 9.1	dbj BAA09767 .1
HCROO13R (AF164515) Cps7G [Streptococcus suis] >sp AAF18954 AAF18954 Cps7G. Length = 404	HCQDW65R (AK001659) unnamed protein product [Homo sapiens] Length = 359	Ag immunoglobulin Fab kapp: Length = 214	91318.1  TBP- tpiens] or [Homo ption factor	ribosomal protein L3 [Homo sapiens] >emb CAA18450.1  (AL022326) dJ333H23.1 (60S Ribosomal Protein L3) [Homo sapiens] >pir S34195 S34195 ribosomal protein L3, cytosolic - human >sp P39023 RL3_HUMAN 60S RIBOSOMAL PROTEIN L3 (HIV-1 TAR RNA BINDING PROTEIN B)		nsin -		H2CAA07R (AJ012449) NSI-binding protein [Homo sapiens] >sp Q9Y480 Q9Y480 NSI-BINDING PROTEIN. Length = 619	The KIAA0146 gene product is novel. [Homo sapiens] >sp Q14159 Q14159 KIAA0146 PROTEIN (FRAGMENT). Length = 918
HCR0013R	HCQDW65R	HCQCV70R	HCQDN27R	HCQCI92R	HCRMP36RA	HCROV67R	HCROT79R	H2CAA07R	H2LAD20R
2638	2639	2640	2641	2642	2643	2644	2645	2646	2647

pSport1	pCMVSport 2.0	pSport1	pSport1	pSport1	Lambda ZAP II	pBluescript	pSport1
HWLQZ32	HKAOU89	HWLU092	HCROW19	HCRQK79	HCQAD53	HKCUD58	HCRNR93
61	70	97	96	82		81	82
61	20	97	96	82		08	82
405	280	112	147	258	96	323	334
28	29	2	-	-	257	45	2
6925	9269	6927	6928	6929	6930	6931	6932
gb AAB04939.	gb AAA61151. 1	emb CAA0143	emb CAA3738	gb AAA61154. 1		emb CAA5048 4.1	dbj BAA07598 .1
HWLQZ32R [threonyl-tRNA synthetase [Homo sapiens] >pir A38867 YSHUT threoninetRNA ligase (EC 6.1.1.3) - human >sp P26639 SYTC_HUMAN THREONYL-TRNA SYNTHETASE, CYTOPLASMIC (EC 6.1.1.3) (THREONINE-TRNA LIGASE) (THRRS). Length = 712	HKAOU89R tissue factor precursor [Homo sapiens] Length = 295	HWLUO92R tissue-specific secretory protein [synthetic construct] emb CAA0143   2emb CAA47928.1  orf [Homo sapiens]   1.1    2eb AA67077.1  epididymal secretory protein precursor [Pan troglodytes] >emb CAA55013.1  epididymal secretory protein 14.6 [Macaca fascicularis] >pir[153929][5392	HNF-1 peptides [Rattus norvegicus] Length = 464	transcription factor Sp-1 [Homo sapiens] >pir[A29635]A29635 transcription factor Sp1 - human (fragment) >sp P08047 SP1_HUMAN TRANSCRIPTION FACTOR SP1 (FRAGMENT). Length = 696		HKCUD58R trypsinogen IV b-form [Homo sapiens] emb CAA5048 >sp Q15665 Q15665 TRYPSINOGEN IV B-FORM. 4.1  Length = 259	TSC-22 [Homo sapiens] >emb CAA10951.1  (AJ222700) TSC-22 [Homo sapiens] >gb AAC50566.1  TSC-22 protein [Homo sapiens] >pir JC4813 JC4813 TGF beta-stimulated clone-22 protein - human >sp Q15714 TS22_HUMAN PUTATIVE REGULATORY PROTEIN TSC-22 (TGFB STIMULATED
HWLQZ32R	HKAOU89R	HWLU092R	HCROW19R	HCRQK79R	HCQAD53R	HKCUD58R	HCRNR93R
2648	2649	2650	2651	2652	2653	2654	2655

pSport1	pBluescript SK-	pBluescript SK-	pSport1	pSport1	pBluescript SK-	pBluescript SK-	pBluescript SK-
нwlqніз	H2CBQ60	H2LAW43	HWLVJ22	HWLRQ77	H2CAA28	H2CAA36	H2CBF10
100	86	93	100	66			
100	86	88	100	86			
487	495	584	259	596	256	109	352
2	211	6	2	r.	7	2	209
6933	6934	6935	6936	6937	6938	6639	6940
gb AAA36528. 1	dbj BAA08091 .1	emb CAA7672 0.1	dbjlBAA83996 .1	emb CAA5938			
HWLQH13R protein tyrosine phosphatase (EC 3.1.3.48) [Homo sapiens] >pir A36065 A36065 protein-tyrosine-phosphatase (EC 3.1.3.48), receptor type alpha precursor - human Length = 802	'me 02181	ubiquitin-conjugating enzyme [Mus musculus] >pir[T31067 T31067 BIR repeat containing ubiquitin-conjugating enzyme BRUCE - mouse >sp O88738 O88738 UBIQUITIN-CONJUGATING ENZYME. Length = 4845	(AB032025) ubiquitin [Canis familiaris] -dbj BAA89414.1  (AB036698) ubiquitin [Felis catus] >gb AAB52914.1  ubiquitin/ribosomal fusion protein [Sus scrofa] >emb CAA40313.1  ubiquitin-52 amino acid fusion protein [Homo sapiens] >emb CAA40312.1  ubiquitin-5	UDP-GalNAc:polypeptide N-acetylgalactosaminyl transferase [Homo sapiens] >pirl[137405 137405 polypeptide N-acetylgalactosaminyltransferase (EC 2.4.1.41) - human >sp Q10471 Q10471 POLYPEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE (EC 2.4.1.41) (PROTEIN-UDP ACET			
нwLQн13R	H2CBQ60R	H2LAW43R	HWLVJ22R	HWLRQ77R	H2CAA28R	H2CAA36R	H2CBF10R
2656	2657	2658	2659	2660	2661	2992	2663

pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pSport1									
H2CBG84	H2CBJ35	H2CBJ62	H2CBK71	H2CBN87	H2CBP73	H2CBS94	H2CBV19	H2CBV81	H2CBW73	H2LAD79	H2LAJ28	H2LAZ29	H2LAZ92	H2LBA33	H2LBB20	HAAAJ56
									100						100	
									100						100	
315	366	252	466	310	282	172	413	185	365	231	370	409	355	326	343	78
157		94	275	176	28	2	189	6	84	13	23	134	71	108	2	-
6941	6942	6943	6944	6945	6946	6947	6948	6949	0569	6951	6952	6953	6954	9559	9569	6957
									gb AAD46135.						dbj BAA86440  -1	
									H2CBW73RB (AF080171) zinc finger protein ZNF232 [Homo sapiens] >sp AAD46135 AAD46135 Zinc finger protein ZNF232. Length = 417						(AB032952) KIAA1126 protein [Homo sapiens] >sp BAA86440 BAA86440 KIAA1126 protein (fragment). Length = 618	
H2CBG84R	H2CBJ35R	H2CBJ62R	H2CBK71R	H2CBN87R	H2CBP73R	H2CBS94R	H2CBV19R	H2CBV81R	H2CBW73RB	H2LAD79R	H2LAJ28RB	H2LAZ29R	H2LAZ92R	H2LBA33R	H2LBB20R	HAAAJ56R
2664	2665	2666	2667	2668	2669	2670	2671	2672	2673	2674	2675	2676	2677	2678	2679	2680

pBluescript	Uni-ZAP XR	pSport1	pBluescript	Uni-ZAP XR	pSport1	pSport1		Lambda ZAP II	HCQAB43 Lambda ZAP	Lambda ZAP II	Lambda ZAP II	HCQAC03 Lambda ZAP	HCQAC24 Lambda ZAP	HCQAC80 Lambda ZAP	HCQADI9 Lambda ZAP	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II
HADTN09	HAUBK53	HBAHC91	HBMCP86	HCEOM04	HCFOE14	HCHOX67	HCQAB27	HCQAB42	НСОАВ43	HCQAB44	HCQAB53	HCQAC03	HCQAC24	HCQAC80	HCQAD19	HCQAD25	HCQAD31	нсоар62	HCQAD71
	×																		
87	219	147	280	393	320	287	240	93	422	474	285	345	437	179	197	138	158	319	485
4	1	1	62	142	111	99	127	1	186	130	26	49	228	3	21	1	3	89	291
8569	6569	0969	6961	6962	6963	6964	5969	9969	<i>L</i> 969	8969	6969	0269	6971	6972	6973	6974	5269	9269	2269
HADTN09R	HAUBK53R	HBAHC91R	HBMCP86R	HCEOM04R	HCF0E14R	HCHOX67R	HCQAB27R	HCQAB42R	HCQAB43R	НСQАВ44R	HCQAB53R	HCQAC03R	HCQAC24R	HCQAC80R	нсQAD19R	HCQAD25R	HCQAD31R	HCQAD62R	HCQAD71R
2681	2682	2683	2684	2685	2686	2687	2688	2689	2690	2691	2692	2693	2694	2695	2696	2697	2698	5696	2700

HCQAE24 Lambda ZAP	HCQAE30 Lambda ZAP	HCQAE32 Lambda ZAP	HCQAE39 Lambda ZAP	HCQAF13 Lambda ZAP	HCQAF78 Lambda ZAP	HCQAF94 Lambda ZAP	HCQAG17 Lambda ZAP	HCQAG32 Lambda ZAP	HCQAG34 Lambda ZAP	HCQAG93 Lambda ZAP	HCQAH27 Lambda ZAP	HCQAH33 Lambda ZAP	HCQAH54 Lambda ZAP	HCQAH63 Lambda ZAP	HCQAH89 Lambda ZAP	HCQAII5 Lambda ZAP	HCQAJ29 Lambda ZAP
312 491	304 498	234 437	3 260	125 373	17 394	326 499	26 178	243 494	3 398	2 193	11 100	204 494	3 131	324 494	190 489	1 207	99 287
6978 3	6979 3	6980 2	6981	6982 1	6983	6984 3	6985	6986 2	2869	8869	6869	6990	6991	6992 3	6993   1	6994	5669
HCQAE24R	HCQAE30R	HCQAE32R	HCQAE39R	HCQAF13R	HCQAF78R	HCQAF94R	HCQAG17R	HCQAG32R	HCQAG34R	HCQAG93R	HCQAH27R	HCQAH33R	НСQАН54R	НСОАН63К	НСQАН89В	HCQAII5R	HCQAJ29R
2701	2702	2703	2704	2705	2706	2707	2708	2709	2710	2711	2712	2713	2714	2715	2716	2717	2718

Lambda ZAP II	Lambda ZAP II	HCQAK16 Lambda ZAP	Lambda ZAP II	HCQAM57 Lambda ZAP	HCQAM70 Lambda ZAP	HCQAM78 Lambda ZAP	HCQAN20 Lambda ZAP	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II	HCQAN74 Lambda ZAP				
HCQAJ44	НСQАJ49	HCQAK16	HCQAK17	HCQAK38	HCQAL71	HCQAL81	НСОАМ32	HCQAM57	HCQAM70	HCQAM78	HCQAN20	HCQAN43	HCQAN44	HCQAN53	HCQAN74
63		98													
55		84													
320	263	345	241	300	299	300	203	334	384	247	306	358	342	258	345
<sub>6</sub>	45	-	2	70	201	139	e.	170	175	62	181	134	166	1	-
9669	2669	8669	6669	7000	7001	7002	7003	7004	7005	9002	7007	7008	7009	7010	7011
gb AAF29011. 1 AF1614		gb AAD55678. 1 AF1477													
HCQAJ44R (AF161451) HSPC333 [Homo sapiens] >sp AAF29011 AAF29011 HSPC333 (fragment). Length = 147		HCQAK16R (AF147790) transmembrane mucin 12 [Homo gb AAD55 sapiens] > sp AAD55678 AAD55678 Transmembrane 1 AF1477 mucin 12 (fragment). Length = 585												·	
HCQAJ44R	HCQAJ49R	HCQAK16R	HCQAK17R	HCQAK38R	HCQAL71R	HCQAL81R	HCQAM32R	HCQAM57R	HCQAM70R	HCQAM78R	HCQAN20R	HCQAN43R	HCQAN44R	<b>H</b> CQAN53R	HCQAN74R
2719	2720	2721	2722	2723	2724	2725	2726	2727	2728	2729	2730	2731	2732	2733	2734

HCQAN95 Lambda ZAP	HCQAQ35 Lambda ZAP	HCQAQ94 Lambda ZAP	Lambda ZAP II	Lambda ZAP II	HCQAR70 Lambda ZAP	HCQAR86 Lambda ZAP	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II		Lambda ZAP II					
HCQAN95	<b>НС</b> QАQ35	НСОАО94	HCQAR19	HCQAR63	HCQAR70	HCQAR86	HCQAS25	HCQAS32	HCQAS58	HCQAS60	HCQAS89	HCQAT10	HCQAT12	HCQAT52	HCQAT57	HCQAT94	HCQAV18
				∞	5	5	=+		10	4	.1		7	0	∞	4	S
208 501	3 74	2 82	1 54	271 468	258 395	307 405	1 84	1 93	1 75	3 224	230 361	1 51	56 157	128   280	97 258	136 234	2 235
7012 2	7013	7014	7015	7016 2	7017 · 2	7018 3	7019	7020	7021	7022		7024		7026   1	7027	7028 1	7029
	,									<u> </u>							
												i					
:																	
						;						,			1		
HCQAN95R	HCQAQ35R	HCQAQ94R	HCQAR19R	HCQAR63R	HCQAR70R	HCQAR86R	HCQAS25R	HCQAS32R	HCQAS58R	HCQAS60R	HCQAS89R	HCQAT10R	HCQAT12R	HCQAT52R	HCQAT57R	HCQAT94R	HCQAV18R
2735	2736	2737	2738	2739	2740	2741	2742	2743	2744	2745	2746	2747	2748	2749	2750	2751	2752

HCQAV23 Lambda ZAP II	758 Lambda ZAP II	HCQAV66 Lambda ZAP	HCQAV73 Lambda ZAP	HCQAW23 Lambda ZAP	HCQAW26 Lambda ZAP	HCQAW40 Lambda ZAP	HCQAW67 Lambda ZAP	HCQBA47 Lambda ZAP	\cdot 89 Lambda ZAP II	101 Lambda ZAP II		E13 Lambda ZAP II	E19 Lambda ZAP II			I24 Lambda ZAP II	I79   Lambda ZAP II
HCQAV	HCQAV58	HCQAV	HCQAV	HCQAV	HCQAV	HCQAV	HCQAV	HCQB₽	нсова89	нсовр01	нсове07	нсове13	нсове19	нсовезз	нсовезя	нсовн24	НСОВН79
255	77	96	148	51	70	227	566	144	381	253	51	115	288	166	403	285	261
55	.8	-	2		2	8	3	1	85	65	1	2	82	2	266	106	46
7030	7031	7032	7033	7034	7035	7036	7037	7038	7039	7040	7041	7042	7043	7044	7045	7046	7047
HCQAV23R	HCQAV58R	HCQAV66R	HCQAV73R	HCQAW23R	HCQAW26R	HCQAW40R	HCQAW67R	HCQBA47R	нсова89к	HCQBD01R	нсове07к	HCQBE13R	нсове19к	нсове53к	нсовезяк	нсовн24к	нсовн79к
2753	2754	2755	2756	2757	2758	2759	2760	2761	2762	2763	2764	2765	2766	2767	2768	2769	2770

HCQCB91 Lambda ZAP	HCQCC50 Lambda ZAP	HCQCC51 Lambda ZAP	HCQCC72 Lambda ZAP	HCQCC88 Lambda ZAP	HCQCC93 Lambda ZAP	HCQCD10 Lambda ZAP	HCQCD46 Lambda ZAP	HCQCE19 Lambda ZAP	HCQCE22 Lambda ZAP	HCQCE28 Lambda ZAP	HCQCE32 Lambda ZAP	HCQCE42 Lambda ZAP	HCQCE46 Lambda ZAP	HCQCE59 Lambda ZAP	HCQCE68 Lambda ZAP	HCQCE72 Lambda ZAP	HCQCE79 Lambda ZAP
HC	HC	ЭН	H	ЭН	ЭН	ЭН	ЭН	)H	ЭН	ЭН	ЭН	)H	ЭН	ЭН	ЭН	ЭН	H
66	115	115	416	196	240	437	384	447	446	123	192	103	199	231	427	453	139
1	2	2	192	35	139	84	211	43	93		-	2	2		263		2
7066	1901	8902	6902	7070	7071	7072	7073	7074	7075	9/0/	7077	7078	7079	7080	7081	7082	7083
						PARAMETER C. B. B. C.						:					
HCQCB91R	HCQCC50R	HCQCC51R	HCQCC72R	HCQCC88R	<b>НС</b> QСС93R	HCQCD10R	HCQCD46R	HCQCE19R	HCQCE22R	HCQCE28R	HCQCE32R	HCQCE42R	HCQCE46R	HCQCE59R	HCQCE68R	HCQCE72R	нсосе79к
2789	2790	2791	2792	2793	2794	2795	2796	2797	2798	2799	2800	2801	2802	2803	2804	2805	2806

HCQCE80 Lambda ZAP	HCQCE83 Lambda ZAP	HCQCE92 Lambda ZAP	HCQCE95 Lambda ZAP	HCQCE96 Lambda ZAP	HCQCF26 Lambda ZAP	HCQCF39 Lambda ZAP	HCQCF74 Lambda ZAP	HCQCF77 Lambda ZAP	HCQCF80 Lambda ZAP	HCQCF82 Lambda ZAP	HCQCF88 Lambda ZAP	HCQCG14 Lambda ZAP	HCQCG19   Lambda ZAP II	HCQCG42 Lambda ZAP	HCQCG49   Lambda ZAP	HCQCG56 Lambda ZAP	HCQCG74 Lambda ZAP
HC		HCC				HC	OH									HCC	HC
76	139	75	349	8 437	224	52	76	1 506	117	7 269	168	111	123	105	130	66	69
2	2		2	48	c.	2	2	261	<b>-</b>	57		1			2	1	1
7084	7085	7086	7087	7088	7089	7090	7091	7092	7093	7094	7095	9602	7097	7098	4060	7100	7101
HCQCE80R	HCQCE83R	HCQCE92R	нсосе95R	нсосе 96К	HCQCF26R	HCQCF39R	HCQCF74R	HCQCF77R	HCQCF80R	HCQCF82R	HCQCF88R	HCQCG14R	HCQCG19R	HCQCG42R	HCQCG49R	HCQCG56R	HCQCG74R
2807	2808	2809	2810	2811	2812	2813	2814	2815	2816	2817	2818	2819	2820	2821	2822	2823	2824

HCQCH01 Lambda ZAP	HCQCH03 Lambda ZAP	HCQCH16 Lambda ZAP	HCQCH30 Lambda ZAP	Lambda ZAP II	HCQCH33 Lambda ZAP	HCQCH47   Lambda ZAP	HCQCH61 Lambda ZAP	HCQCH69 Lambda ZAP	HCQCH83 Lambda ZAP	Lambda ZAP II	HCQCI75 Lambda ZAP II						
нсосно1	нсосн03	нсосн16	нсосн30	нсосн32	нсоснзз	нсосн47	нсосн61	нсосне9	нсосн83				HCQCI57	нсост63	HCQCI64	69ГОООН	нсост2
82	295	235	111	282	100	239	83	129	86	309	102	231	96	213	302	96	09
2	41	26		-	2	3 2	3	-	3	103	1	1	-	. 92	120	1	_
7102	7103	7104	7105	7106	7107	7108	7109	7110	7111	7112	7113	7114	7115	7116	7117	7118	7119
									_								
	i i																
HCQCH01R	нсосн03к	нсосн168	нсосн30к	<b>НС</b> QСН32R	нсоснззк	HCQCH47R	нсосныя	нсосн698	<b>НС</b> QСН83R	HCQCI18R	HCQCI28R	HCQCI42R	HCQCI57R	HCQCI63R	HCQCI64R	HCQCI69R	HCQCI75R
2825	2826	2827	2828	2829	2830	2831	2832	2833	2834	2835	2836	2837	2838	2839	2840	2841	2842

2843	HCQCI89R	7120	166	360	нсосів9	Lambda ZAP II
2844	HCQCJIIR	7121	3	176		Lambda ZAP II
2845	HCQCJ21R	7122	-	111	НСОСЛ21	Lambda ZAP II
2846	HCQC125R	7123	09	173	нсосл25	Lambda ZAP II
2847	HCQCJ34R	7124	-	66	нсослз4	HCQCJ34 Lambda ZAP
2848	HCQCJ38R	7125	41	313	нсослз8	HCQCJ38 Lambda ZAP
2849	HCQCJ42R	7126	43	234	НСОСЈ42	Lambda ZAP II
2850	HCQCJ45R	7127	7	75	нсосл45	Lambda ZAP II
2851	HCQCJ50R	7128	1	66	нсослзо	Lambda ZAP II
2852	HCQCJ51R	7129	3	116	нсосля	Lambda ZAP II
2853	HCQCJ68R	7130	1	66		Lambda ZAP II
2854	HCQCJ76R	7131	1	48		Lambda ZAP II
2855	HCQCJ77R	7132	2	130	нсосл77	Lambda ZAP II
2856	HCQCJ85R	7133	1	129	HCQCJ85	Lambda ZAP II
2857	HCQCJ89R	7134	182	415		Lambda ZAP II
2858	HCQCJ94R	7135	46	207	НСОСЛ94	Lambda ZAP II
2859	HCQCK03R	7136	2	103	нсоск03	Lambda ZAP II
2860	HCQCK17R	7137	-	246	нсоск17	HCQCK17 Lambda ZAP

HCQCK25 Lambda ZAP	HCQCK34 Lambda ZAP	HCQCK39 Lambda ZAP	HCQCK50 Lambda ZAP	HCQCK54 Lambda ZAP	HCQCK58 Lambda ZAP	HCQCK59 Lambda ZAP	HCQCK81 Lambda ZAP	HCQCK90 Lambda ZAP	HCQCL01 Lambda ZAP	HCQCL05 Lambda ZAP	HCQCL07 Lambda ZAP	HCQCL11 Lambda ZAP	HCQCL14 Lambda ZAP	HCQCL19 Lambda ZAP	HCQCL20 Lambda ZAP	HCQCL22 Lambda ZAP	HCQCL30 Lambda ZAP
	<u> </u>	Н	<u> </u>	<u> </u>		<u></u>		I	-								
159	114	140	103	66	121	280	380	201	445	252	89	102	229	104	187	279	101
-	1	3	2	-	2	119	153	22	143	91	3	1	2	т	∞	142	27
7138	7139	7140	7141	7142	7143	7144	7145	7146	7147	7148	7149	7150	7151	7152	7153	7154	7155
HCQCK25R	HCQCK34R	нсоск39к	HCQCK50R	HCQCK54R	HCQCK58R	HCQCK59R	HCQCK81R	HCQCK90R	HCQCL01R	HCQCL05R	нсостоля	HCQCL11R	HCQCL14R	HCQCL19R	HCQCL20R	HCQCL22R	HCQCL30R
2861	2862	2863	2864	2865	2866	2867	2868	2869	2870	2871	2872	2873	2874	2875	2876	2877	2878

HCQCL3SR         7156         1         102           HCQCL4RR         7157         1         237           HCQCL4RR         7158         1         69           HCQCL4RR         7159         57         251           HCQCL5RR         7160         2         100           HCQCL5RR         7161         17         133           HCQCL5RR         7163         2         253           HCQCL6RR         7164         1         144           HCQCL6RR         7166         1         211           HCQCL6RR         7166         1         214           HCQCL73R         7167         3         155           HCQCL73R         7169         151         330           HCQCL73R         7170         34         177           HCQCL90R         7171         3         278           HCQCL90R         7171         3         278           HCQCL90R         7172         6         134           HCQCMG0R         7173         7173         387	HCQCL35 Lambda ZAP	HCQCL43 Lambda ZAP	HCQCL46 Lambda ZAP	HCQCL48 Lambda ZAP	HCQCL51 Lambda ZAP	HCQCL54 Lambda ZAP	HCQCL55 Lambda ZAP	HCQCL63 Lambda ZAP	HCQCL64 Lambda ZAP	HCQCL65 Lambda ZAP	HCQCL66 Lambda ZAP		HCQCL73 Lambda ZAP	HCQCL78 Lambda ZAP	HCQCL79 Lambda ZAP	HCQCL90 Lambda ZAP	HCQCL92 Lambda ZAP	HCQCM69 Lambda ZAP
HCQCL3SR         7156           HCQCL4SR         7158           HCQCL4RR         7160           HCQCL5RR         7161           HCQCL5RR         7161           HCQCL5RR         7165           HCQCL6RR         7165           HCQCL6RR         7166           HCQCL6RR         7166           HCQCL5RR         7167           HCQCL7SR         7167           HCQCL7SR         7169           HCQCL7SR         7170           HCQCL9RR         7170           HCQCL9RR         7171           HCQCL9RR         7171           HCQCL9RR         7171           HCQCL9RR         7171	102	237	69	251	100	133	100	253	144	280	231	155	214	330	177	278	134	387
HCQCL35R HCQCL43R HCQCL48R HCQCL51R HCQCL51R HCQCL53R HCQCL63R HCQCL73R HCQCL73R HCQCL73R HCQCL73R HCQCL73R	-	-	-	57	2	17	2	2	-	2	1	3	110	151	34	3	9	172
	7156	7157	7158	7159	7160	7161	7162	7163	7164	7165	7166	7167	7168	7169	7170	7171	7172	7173
2879       2880       2881       2882       2883       2884       2889       2889       2889       2889       2890       2891       2892       2893       2893       2894       2895       2896       2897       2898       2899       2899       2890 <td>2879 HCQCL35R</td> <td>2880 HCQCL43R</td> <td>-</td> <td>2882 HCQCL48R</td> <td>2883 HCQCL51R</td> <td>2884 HCQCL54R</td> <td>2885 HCQCL55R</td> <td>2886 HCQCL63R</td> <td>2887 HCQCL64R</td> <td>2888 HCQCL65R</td> <td>2889 HCQCL66R</td> <td>2890 HCQCL69R</td> <td>2891 HCQCL/3R</td> <td>2892 HCQCL78R</td> <td>2893 HCQCL79R</td> <td>2894 HCQCL90R</td> <td>2895 HCQCL92R</td> <td>2896 HCQCM69R</td>	2879 HCQCL35R	2880 HCQCL43R	-	2882 HCQCL48R	2883 HCQCL51R	2884 HCQCL54R	2885 HCQCL55R	2886 HCQCL63R	2887 HCQCL64R	2888 HCQCL65R	2889 HCQCL66R	2890 HCQCL69R	2891 HCQCL/3R	2892 HCQCL78R	2893 HCQCL79R	2894 HCQCL90R	2895 HCQCL92R	2896 HCQCM69R

7174 2 193 HCQCO30 Lambda ZAP	7175 122 388 HCQCO53 Lambda ZAP	7176 82 273 HCQCO57 Lambda ZAP	7177 1 162 HCQCO66 Lambda ZAP	7178 29 151 HCQCO79 Lambda ZAP	7179 2 139 HCQCO85 Lambda ZAP II	7180 3 338 HCQCP08 Lambda ZAP	7181 108 296 HCQCP14 Lambda ZAP	7182 2 55 HCQCP15 Lambda ZAP II	7183 15 128 HCQCP19 Lambda ZAP	7184 1 105 HCQCP23 Lambda ZAP	7185 213 365 HCQCP27 Lambda ZAP	7186 102 200 HCQCP30 Lambda ZAP	7187 3 62 HCOCP35 Lambda ZAP		1 48	1 48	1 48
	нсособзя	нсосозля	нсособбя	нсосо79к	нсосовяя	HCQCP08R	НСQСР14R	нсосрія	нсоср19к	нсоср23К	нсоср27к	нсосрзок	HCQCP35R		HCQCP42R	HCQCP42R HCQCP58R	HCQCP42R HCQCP58R HCQCP75R
2897 HCQCO30R	2898 H	2899 Н	2900 Н	2901 Н	2902 Н	2903 Н	2904 H	2905 Н	2906 Н	2907 H	2908 Н	2909 Н	2910 Н	2011 I			

HCQCP86 Lambda ZAP	HCQCP89 Lambda ZAP	HCQCQ09 Lambda ZAP	HCQCQ17 Lambda ZAP	HCQCQ48 Lambda ZAP	HCQCR15 Lambda ZAP	HCQCR44 Lambda ZAP	HCQCR69 Lambda ZAP	HCQCT38 Lambda ZAP	HCQCT49 Lambda ZAP	HCQCT84 Lambda ZAP	HCQCT89 Lambda ZAP	HCQCU08 Lambda ZAP	HCQCU19 Lambda ZAP	HCQCU37 Lambda ZAP	HCQCUSS Lambda ZAP	HCQCU57 Lambda ZAP	HCQCU59 Lambda ZAP
НС	НС	HC	HC	HC HC	НС	НС	ЭН	НС	ЭН	ЭН	ЭН НС	HC	ЭН	ОН	ЭН	он Н	HC
66	166	575	159	111	436	400	186	184	158	198	367	122	453	97	486	456	51
1	2	300	22	-	140	173	70	71	3	1	2	3	235	. 2	265	274	-
7192	7193	7194	7195	7196	7197	7198	7199	7200	7201	7202	7203	7204	7205	7206	7207	7208	7209
нсосрявя	нсосрвяя	нсосооя не	нсосо17R	HCQCQ48R	HCQCR15R	HCQCR44R	нсоск69к	нсостзяя	нсост49к	HCQCT84R	нсоствяя	HCQCU08R	HCQCU19R	HCQCU37R	нсосизя	HCQCU57R	нсосизэк
2915	2916	2917	2918	2919	2920	2921	2922	2923	2924	2925	2926	2927	2928	2929	2930	2931	2932

HCQCU67 Lambda ZAP	HCQCU72 Lambda ZAP	HCQCU73 Lambda ZAP	HCQCV01 Lambda ZAP	HCQCV21 Lambda ZAP	HCQCV50 Lambda ZAP	HCQCV68 Lambda ZAP	HCQCV73 Lambda ZAP	HCQCV91 Lambda ZAP	HCQCX11 Lambda ZAP	Lambda ZAP II	Lambda ZAP II	Lambda ZAP II	HCQCX33 Lambda ZAP	Lambda ZAP II	HCQCX90 Lambda ZAP	HCQDA09 Lambda ZAP	HCQDA20 Lambda ZAP
НСОСО67	нсоси72	нсоси73	НСОСУ01	нсосу21	нсосу50	НСОСУ68	НСОСУ73	НСОСУ91	HCQCX11	HCQCX18	нсосх21	нсосх22	нсосх33	нсосх <i>51</i>	нсосх90	НСОДА09	HCQDA20
168	238	66	628	66	584	625	183	227	524	264	262	364	277	399	344	989	. 704
-	131	-	371	-	330	425		84	177	100	209	2	32	166	213	346	375
7210	7211	7212	7213	7214	7215	7216	7217	7218	7219	7220	7221	7222	7223	7224	7225	7226	7227
R	R	R	R	<b>8</b>	R	8	R	R	R	R	<b>8</b>	R	<b>X</b>	R	R	~	Ж
HCQCU67R	HCQCU72R	HCQCU73R	HCQCV01R	HCQCV21R	HCQCV50R	HCQCV68R	HCQCV73R	HCQCV91R	HCQCX11R	HCQCX18R	HCQCX21R	HCQCX22R	HCQCX33R	HCQCX57R	HCQCX90R	HCQDA09R	HCQDA20R
2933	2934	2935	2936	2937	2938	2939	2940	2941	2942	2943	2944	2945	2946	2947	2948	2949	2950

HCQDA28 Lambda ZAP	HCQDA36 Lambda ZAP	HCQDA52 Lambda ZAP	HCQDA55 Lambda ZAP	HCQDA66 Lambda ZAP	HCQDA86 Lambda ZAP	7 Lambda ZAP II	6 Lambda ZAP II	HCQDB29 Lambda ZAP	HCQDB41 Lambda ZAP	8 Lambda ZAP II	HCQDB49 Lambda ZAP	2 Lambda ZAP II	4 Lambda ZAP II	5 Lambda ZAP II	8 Lambda ZAP II	2 Lambda ZAP II	2 Lambda ZAP II
HCQDA2	нсораз	HCQDAS;	НСОДА	НСОДА6	HCQDA8	нсорв17	нсорв26	нсорва	нсорв4	HCQDB48	нсорв4	нсорв52	нсорвя	НСОВБ	нсорв78	нсорс05	нсорс15
349	141	102	454	252	333	268	494	148	150	461	120	373	391	293	09	185	329
89		-	176	-	238	2	303	2	61	282	1	2	2	126	1	3	156
7228	7229	7230	7231	7232	7233	7234	7235	7236	7237	7238	7239	7240	7241	7242	7243	7244	7245
HCQDA28R	HCQDA36R	HCQDA52R	HCQDA55R	HCQDA66R	HCQDA86R	HCQDB17R	нсфрв26К	нсфрв29к	HCQDB41R	HCQDB48R	НСQDВ49R	нсфрв52R	НСQDB54R	нсфрв55R	HCQDB78R	нсQDC02R	нсфрс12R
2951	2952	2953	2954	2955	2956	2957	2958	2959	2960	2961	2962	2963	2964	2965	2966	2967	2968

HCQDC13 Lambda ZAP	HCQDC15 Lambda ZAP	HCQDC28 Lambda ZAP	HCQDC29 Lambda ZAP	Lambda ZAP II	HCQDC44 Lambda ZAP	Lambda ZAP II	HCQDC74 Lambda ZAP	HCQDC88 Lambda ZAP	HCQDD35 Lambda ZAP	Lambda ZAP II							
нсорс13	HCQDC15	нсорс28	нсорс29	нсорс33	HCQDC44	нсорсез	HCQDC74	HCQDC88	нсоррзя	нсорр65	нсорр91	нсорео4	нсорето	нсорб20		нсорез1	нсорез8
209	94	155	317	175	542	809	360	515	224	617	465	86	396	66	482	66	322
45	2	45	132	17	408	273	145	303	06	270	316	24	163	1	255	1	107
7246	7247	7248	7249	7250	7251	7252	7253	7254	7255	7256	7257	7258	7259	7260	7261	7262	7263
HCQDC13R	HCQDC15R	HCQDC28R	нсорс298	нсорсззк	HCQDC44R	нсорс638	HCQDC74R	HCQDC88R	нсоррзя	нсорр658	нсоррэтк	HCQDE04R	HCQDE10R	HCQDE20R	<b>НСQDE25</b> R	нсорез18	нсорез8к
2969	2970	2971	2972	2973	2974	2975	2976	2977	2978	2979	2980	2981	2982	2983	2984	2985	2986

HCQDE45 Lambda ZAP	HCQDE52 Lambda ZAP	HCQDE58 Lambda ZAP	HCQDE59 Lambda ZAP	HCQDE61 Lambda ZAP	HCQDE68 Lambda ZAP	HCQDF22 Lambda ZAP	HCQDF44 Lambda ZAP	HCQDF51 Lambda ZAP	HCQDF66 Lambda ZAP	HCQDF69 Lambda ZAP	HCQDF70 Lambda ZAP	HCQDF79 Lambda ZAP	HCQDF93 Lambda ZAP	HCQDG40 Lambda ZAP	HCQDG62 Lambda ZAP	HCQDG71 Lambda ZAP	HCQDG80 Lambda ZAP
340	265	114	214	198	146	61	5 427	127	111	184	116	129	3 332	279	66	92	2 271
7264 185	7265 74	7266 1	7267 17	7268 1	7269 3	7270 2	7271 125	7272 50	7273 1	7274 2	7275 3	7276 1	7277 153	7278 79	7279 1	7280 21	7281 152
									_								
HCQDE45R	<b>НС</b> QDE52R	HCQDE58R	НСQDE59R	HCQDE61R	НСQDE68R	нсорғал	HCQDF44R	HCQDF51R	<b>НСQDF66R</b>	<b>НС</b> QDF69R	HCQDF70R	HCQDF79R	<b>НСQDF93R</b>	HCQDG40R	HCQDG62R	HCQDG71R	HCQDG80R
2987	2988	2989	2990	2991	2992	2993	2994	2995	2996	2997	2998	2999	3000	3001	3002	3003	3004

	Lambda ZAP II	HCQDH41 Lambda ZAP	HCQDH42 Lambda ZAP	HCQDH50 Lambda ZAP	HCQDH57 Lambda ZAP	HCQDH60 Lambda ZAP II	HCQDH65 Lambda ZAP	HCQDH66 Lambda ZAP	HCQDH68 Lambda ZAP	HCQDH78 Lambda ZAP	Lambda ZAP II	5 Lambda ZAP II	Lambda ZAP II				
HCQDG80	нсорн18	нсорн4	нсорн4;	нсорня	нсорня	нсорне	нсорне:	нсорне	9НСОДН (	нсорн7	нсорн79	нсорн95	нсорп н	НСQDI42	нсорі63	нсорі67	нсоргу
108	141	57	292	443	346	124	89	147	102	102	102	109	54	229	105	541	444
7282 1	7283 31	7284 1	7285 83	7286 150	7287 176	7288 2	7289 6	7290 1	7291 1	7292 1	7293 1	7294 2	7295 1	7296 2	7297 1	7298 290	7299 241
нсорд868	<b>НС</b> QDH18R	нсорн418	<b>НС</b> QDH42R	нсорн508	нсорн578	нсорн608	нсорны	нсорн668	нсорн688	нсорн78R	нсорн798	нсорн95R	HCQDI19R	HCQDI42R	нсор163К	HCQDI67R	HCQDI77R
3005	3006	3007	3008	3009	3010	3011	3012	3013	3014	3015	3016	3017	3018	3019	3020	3021	3022

HCQDI78 Lambda ZAP	HCQDI85 Lambda ZAP	HCQDI95 Lambda ZAP	13 Lambda ZAP II	HCQDJ19 Lambda ZAP	21 Lambda ZAP II	22 Lambda ZAP II	52 Lambda ZAP II	58 Lambda ZAP II	70 Lambda ZAP II	83 Lambda ZAP II	84 Lambda ZAP II	91 Lambda ZAP II	93 Lambda ZAP II	95 Lambda ZAP II	HCQDK13 Lambda ZAP	19 Lambda ZAP II	HCQDK20 Lambda ZAP
НСОБІЛ	НСОДІВ	НСОБІЗ	нсорыз	НСОВЛ	нсорл21	нсор122	нсор152	нсорлея	нсорлу	нсорляз	нсорля4	нсоръя	НСОДР93	нсорля	НСОВК	нсорк19	нсорк
140	427	461	108	544	363	102	102	543	69	111	114	182	445	473	422	393	114
3	185	117	_	299	58	-	-	238	-	1	-	8	152	93	186	259	-
7300	7301	7302	7303	7304	7305	7306	7307	7308	7309	7310	7311	7312	7313	7314	7315	7316	7317
W.	~	<b>8</b>	2	X	2	ж.		R	X.	<b>X</b>	2	Ж	R	R	JR	JR.	J.R.
HCQDI78R	нсорівзя	нсор1958	HCQDJ13R	нсорл19К	HCQDJ21R	HCQDJ22R	HCQDJ52R	HCQDJ68R	HCQDJ70R	HCQD183R	HCQDJ84R	HCQDJ91R	нсор1938	нсор1958	нсорк13R	нсорк 19R	HCQDK20R
3023	3024	3025	3026	3027	3028	3029	3030	3031	3032	3033	3034	3035	3036	3037	3038	3039	3040

HCQDK34   Lambda ZAP	HCQDK49 Lambda ZAP	HCQDK50 Lambda ZAP	HCQDK56 Lambda ZAP	HCQDK58 Lambda ZAP	HCQDK75 Lambda ZAP	HCQDK89 Lambda ZAP	HCQDL12 Lambda ZAP	HCQDL24 Lambda ZAP	HCQDL36 Lambda ZAP	HCQDL43 Lambda ZAP	HCQDL52 Lambda ZAP	HCQDL54 Lambda ZAP	HCQDL57 Lambda ZAP	HCQDL93 Lambda ZAP	HCQDL96 Lambda ZAP	HCQDM01 Lambda ZAP	HCQDM17 Lambda ZAP
ЭЭН											ЭЭН			ЭЭН			ЭЭН
3 59	1 105	1 153	2 154	14 127	1 114	154 360	22 246	3 200	1 132	1 114	1 54	2 292	55 174	1 99	1 135	3 107	3 53
7318	7319	7320	7321	7322	7323	7324	7325	7326	7327	7328	7329	7330	7331	7332	7333	7334	7335
НСQDК34R	HCQDK49R	HCQDK50R	HCQDK56R	HCQDK58R	HCQDK75R	НСQDК89R	HCQDL12R	HCQDL24R	HCQDL36R	HCQDL43R	HCQDL52R	HCQDL54R	HCQDL57R	<b>НСQDL93R</b>	HCQDL96R	HCQDM01R	HCQDM17R
3041	3042	3043	3044	3045	3046	3047	3048	3049	3050	3051	3052	3053	3054	3055	3056	3057	3058

HCQDM49 Lambda ZAP	HCQDM55 Lambda ZAP	HCQDM58 Lambda ZAP	HCQDN08 Lambda ZAP	2 Lambda ZAP II	3 Lambda ZAP II	HCQDN78 Lambda ZAP	HCQDO05 Lambda ZAP	HCQDO07 Lambda ZAP	HCQDO25 Lambda ZAP	HCQDO44 Lambda ZAP	0 Lambda ZAP II	HCQDO75 Lambda ZAP	3 Lambda ZAP II	8 Lambda ZAP II	4 Lambda ZAP II	8 Lambda ZAP II	1 Lambda ZAP II
НСОДМ4	НСОВМЯ	нсормя	HCQDNO	HCQDN32	HCQDN33	HCQDN7	нсоро	нсороо	нсорох	нсоро4	нсорое0	нсорол	НСОДО83	НСОДО88	НСОДР14	нсорыя	НСОДР41
146	99	114	213	99	515	105	286	171	413	296	241	63	300	186	443	240	54
3	1	_	<i>L</i> 9	e.	252	-	122	40	138	69	2	1	112	82	237	61	
7336	7337	7338	7339	7340	7341	7342	7343	7344	7345	7346	7347	7348	7349	7350	7351	7352	7353
нсфрм49К	HCQDM55R	HCQDM58R	HCQDN08R	HCQDN32R	HCQDN33R	HCQDN78R	нсфрообя	нсфроотк	нсфрог5к	нсфро44к	нсфробов	нсqр075R	нсоровзк	нсфро88к	HCQDP14R	нсфрр18к	HCQDP41R
3059	3060	3061	3062	3063	3064	3065	3066	3067	3068	3069	3070	3071	3072	3073	3074	3075	3076

1 102	5 247 390 HCQDP91 Lambda ZAP	6 241 360 HCQDQ08 Lambda ZAP II	7 269 427 HCQDQ09 Lambda ZAP	1 102 HCQDQ45	9 62 283 HCQDQ80 Lambda ZAP	286 516	1 99	2 86 292 HCQDR54 Lambda ZAP	3 216 446 HCQDR89 Lambda ZAP II	4 1 300 HCQDS01 Lambda ZAP	5 177 542 HCQDS14 Lambda ZAP	6 3 152 HCQDS29 Lambda ZAP	7 1 105 HCQDS34 Lambda ZAP	8 130 360 HCQDS40 Lambda ZAP II	9 183 416 HCQDS51 Lambda ZAP II	0 348 548 HCQDS53 Lambda ZAP	339 533 HCQDS56
7354	7355	7356	7357	7358	7359	7360	7361	7362	7363	7364	7365	9982	7367	7368	7369	7370	7371
HCQDP50R	<b>НС</b> QDР91R	<b>НС</b> QDQ08R	нсороо9R	HCQDQ45R	<b>НС</b> QDQ80R	HCQDR04R	HCQDR49R	HCQDR54R	HCQDR89R	HCQDS01R	HCQDS14R	HCQDS29R	HCQDS34R	HCQDS40R	HCQDS51R	HCQDS53R	HCQDS56R
3077	3078	3079	3080	3081	3082	3083	3084	3085	3086	3087	3088	3089	3090	3091	3092	3093	3094

	Lambda ZAP II	HCQDT48 Lambda ZAP	Lambda ZAP	HCQDT64 Lambda ZAP	Lambda ZAP II	5 Lambda ZAP II	Lambda ZAP II										
HCQDS61	нсорѕ62	нсорѕез	HCQDS67	нсорѕ83	HCQDS84	HCQDS89	HCQDT17	HCQDT48	нсорт63	нсорт64	нсорт93	HCQDU24	HCQDU34	нсориз9	нсорие0	нсорие5	НСОБО69
300	290	564	114	290	102	78	733	361	61	347	418	102	128	194	102	115	156
	3	388	-	432 5			398	113	2	141	236	31	3	57		2	1
7372	7373	7374	7375	7376	7377	7378	7379	7380	7381	7382	7383	7384	7385	7386	7387	7388	7389
									:								
HCQDS61R	HCQDS62R	HCQDS63R	HCQDS67R	HCQDS83R	HCQDS84R	HCQDS89R	HCQDT17R	HCQDT48RA	нсортезк	HCQDT64R	нсорт93RA	HCQDU24R	нсориз4к	нсори 59к	HCQDU60R	нсориея	нсори69к
3095	3096	3097	3098	3099	3100	3101	3102	3103	3104	3105	3106	3107	3108	3109	3110	3111	3112

HCQDU94 Lambda ZAP	HCQDV27 Lambda ZAP	HCQDV41 Lambda ZAP	HCQDV44 Lambda ZAP	HCQDV49 Lambda ZAP	HCQDV53 Lambda ZAP	HCQDV62 Lambda ZAP	HCQDV63 Lambda ZAP	HCQDV64 Lambda ZAP	HCQDV74 Lambda ZAP	HCQDV76 Lambda ZAP	HCQDV79 Lambda ZAP	HCQDV83   Lambda ZAP	HCQDW01 Lambda ZAP	HCQDW02 Lambda ZAP	HCQDW15 Lambda ZAP	HCQDW30 Lambda ZAP	HCQDW38 Lambda ZAP
92	112	528	274	419	260	66	174	102	123	105	72	238	127	86	26	100	156
2	17	-	2	87	96	43	37				7	26	2	m	2	2	-
7390	7391	7392	7393	7394	7395	7396	7397	7398	7399	7400	7401	7402	7403	7404	7405	7406	7407
3 HCQDU94R		5 HCQDV41R	-	7 HCQDV49R	8 HCQDV53R	9 HCQDV62R	0 HCQDV63R	1 HCQDV64R					6 HCQDW01R	7 HCQDW02R	8 HCQDW15R	9 HCQDW30R	0 HCQDW38R
3113	3114	3115	3116	3117	3118	3119	3120	3121	3122	3123	3124	3125	3126	3127	3128	3129	3130

7409 2 7410 74 7410 74 7411 11 7411 11 7412 30 7413 11 7414 11 7415 5 7415 5 7416 207 7416 207 7418 2 7419 22 7420 108 7421 2 7421 2 7421 36 7421 36 7421 36 7421 36 7421 36 7421 36 7421 36 7422 16 7422 16		Tal Lal La
7410       74         7411       1         7412       30         7413       1         7414       1         7415       5         7416       207         7417       3         7418       2         7419       22         7420       108         7421       2         7422       3         7423       110         7423       110         7424       36         7425       161         7426       1         7427       3         7428       1         7429       148		Lambda ZAP II PSport1
7411       1         7412       30         7413       1         7414       1         7415       5         7416       207         7417       3         7418       2         7419       2         7410       207         7411       3         7411       3         7421       2         7422       3         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148		
7412       30         7413       1         7414       1         7415       5         7416       207         7417       3         7418       2         7420       108         7421       2         7421       2         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148		
7413       1         7414       1         7415       5         7416       207         7417       3         7418       2         7419       22         7420       108         7421       2         7422       3         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148		
7416       1         7415       5         7416       207         7417       3         7418       2         7419       22         7420       108         7421       2         7422       3         7424       36         7426       1         7426       1         7427       2         7428       1         7429       148		
7415       5         7416       207         7417       3         7418       2         7419       22         7420       108         7421       2         7422       3         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148		
7416       207         7417       3         7418       2         7419       22         7420       108         7421       2         7422       3         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148		
7417       3         7418       2         7419       22         7420       108         7421       2         7423       110         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148		
7418       2         7419       22         7420       108         7421       2         7422       3         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148		
7419       22         7420       108         7421       2         7422       3         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148         7430       184		
7420       108         7421       2         7422       3         7423       110         7424       36         7425       161         7426       1         7427       2         7428       1         7429       148         7439       184		
7421     2       7422     3       7423     110       7424     36       7425     161       7426     1       7427     2       7429     148       7429     148		
7422     3       7423     110       7424     36       7425     161       7426     1       7427     2       7427     2       7428     1       7429     148		psporti
7423       110         7424       36         7425       161         7426       1         7426       1         7427       2         7429       1         7429       148	HCKMB82	pSport1
7424       36         7425       161         7426       1         7427       2         7427       2         7429       1         7429       148         7430       184		5 pSport1
7425     161       7426     1       7427     2       7428     1       7429     148       7430     184	HCRMC01	pSport1
7426       1         7427       2         7428       1         7429       148         7430       184	HCRMC13	3 pSport1
7427     2       7428     1       7429     148       7430     184	HCRMC85	5 pSport1
7428     1       7429     148       7430     184		l pSport1
7429 148	HCRMD24	t pSport1
7430 184	HCRMD33	3 pSport1
	HCRMD57	7 pSport1
HCRMD77R 3 185	HCRMD77	7 pSport1
HCRME08R   7432   56   214	HCRME08	s pSport1
HCRME25R 7433 1 162	HCRME25	5 pSport1
HCRME49R 7434 41 274		pSport1
HCRMF03R 7435 1 111	HCRMF03	b pSport1

															Γ	Γ	<u> </u>													
pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1																					
HCRMF07	HCRMF23	HCRMF24	HCRMF33	HCRMF38	HCRMF47	HCRMF67	HCRMF72	HCRMF82	HCRMF84	HCRMF91	HCRMF93	HCRMF94	HCRMG20	HCRMG43	HCRMG80	HCRMH08	HCRMH75	HCRMH83	HCRMH94	HCRMI04	HCRMI33	HCRMI40	HCRMI47	HCRMI60	HCRMJ03	HCRMJ21	HCRMJ54	HCRMJ80	HCRMJ81	HCRMJ84
																													100	
																													100	
201	86	305	106	151	473	151	195	234	213	106	375	402	255	201	466	87	84	196	66	346	144	354	62	337	09	208	06	118	473	58
1	3	75	2	50	261	2	-	16	-	2	91	1	139	1	2	1	4	17	1	197	1	229	3	2	-	194	1	2	207	2
7436	7437	7438	7439	7440	7441	7442	7443	7444	7445	7446	7447	7448	7449	7450	7451	7452	7453	7454	7455	7456	7457	7458	7459	7460	7461	7462	7463	7464	7465	7466
																													dbj BAA88923	
																													(AB023584) reduced expression in cancer [Homo sapiens] >sp BAA88923 BAA88923 Rec protein. Length = 367	
HCRMF07R	HCRMF23R	HCRMF24R	HCRMF33R	HCRMF38R	HCRMF47R	HCRMF67R	HCRMF72R	HCRMF82R	HCRMF84R	HCRMF91R	HCRMF93R	HCRMF94R	HCRMG20R	HCRMG43R	HCRMG80R	HCRMH08R	HCRMH75R	HCRMH83R	HCRMH94R	HCRMI04R	<b>HCRMI33R</b>	HCRMI40R	HCRMI47R	HCRMI60R	HCRMJ03R	HCRMJ21R	HCRMJ54R	HCRMJ80R	HCRMJ81R	HCRMJ84R
3159	3160	3161	3162	3163	3164	3165	3166	3167	3168	3169	3170	3171	3172	3173	3174	3175	3176	3177	3178	3179	3180	3181	3182	3183	3184	3185	3186	3187	3188	3189

HCRMK11 pSport1	HCRMK94 pSport1	HCRMN04 pSport1	HCRMO53 pSport1	HCRMO55 pSport1	HCRMP32 pSport1	HCRMR07 pSport1	HCRMR28 pSport1	HCRMR50 pSport1	HCRMR51 pSport1	HCRMS48 pSport1	HCRMS54 pSport1	HCRMS55 pSport1	HCRMT03 pSport1	HCRMT32 pSport1	HCRMU10 pSport1	HCRMU21 pSport1	HCRMU34 pSport1	HCRMU36 pSport1	HCRMU63 pSport1	HCRMU67 pSport1	HCRMU76 pSport1	HCRMU78 pSport1	HCRMU85 pSport1	HCRMV06 pSport1	HCRMV52 pSport1	HCRMV67 pSport1	HCRMV78 pSport1	HCRMV95 pSport1	HCRMW15 pSport1	HCRMW62 pSport1	HCRMW90 pSport1	HCRMX02 pSport1
3 299	30 92	16 201	1 99	3 140	160 390	2 67	2 274	1 144	2 100	159 332	1 99	13 72	2 82	1 69 1	237 476	217 378	130 321	9 131	2 64	1 60	41 79	1 66	3 245	2 82	19 90	2 247	12 146	1 54	2 205	2 283	2 142	1 180
7467	7468	7469	7470	7471	7472	7473	7474	7475	7476		7478	7479	7480	7481	7482	7483		7485	7486	7487	7488	7489	7490	7491	7492	7493	7494	7495	7496	7497	7498	7499
																													-			
HCRMK11R	HCRMK94R	HCRMN04R	HCRM053R	HCRMOSSR	HCRMP32RA	HCRMR07R	HCRMR28R	HCRMR50R	HCRMR51R	HCRMS48R	HCRMS54R	HCRMS55R	HCRMT03R	HCRMT32R	HCRMU10R	HCRMU21R	HCRMU34R	HCRMU36R	HCRMU63R	HCRMU67R	HCRMU76R	HCRMU78R	HCRMU85R	HCRMV06R	HCRMV52R	HCRMV67R	HCRMV78R	HCRMV95R	HCRMW15R	HCRMW62R	HCRMW90R	HCRMX02R
3190 HC	3191 HC	3192 HG	3193 HC	3194 H(	3195 HC	3196 HG	3197 H(	3198 H(	3199 H(	3200 H(	3201 H(	3202 H(	3203 H(	3204 H(	3205 HG	3206 H(	3207 HG	3208 HG	3209 H(	3210 HG	3211 H(	3212 HG	3213 HG	3214 H(	3215 HG	3216 H(	3217 H(	3218 H(	3219 HC		3221 H(	3222 H(

pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1							
HCRMX11	HCRMY28	HCRMY29	HCRMY39	HCRMZ13	HCRMZ36	HCRMZ53	HCRMZ71	HCRMZ92	HCRNA39	HCRNA44	HCRNA64	HCRNA88	HCRNB36	HCRNB47	HCRNB56	HCRNB61	HCRNB69	HCRNB77	HCRNB85	HCRNC23	HCRND21	HCRND28	HCRND30	HCRND45	HCRNE04	HCRNE11	HCRNE15	HCRNE17	HCRNE18
							68														-	·							
502	105	109	157	69	451	16	440 85	482	56	168	108	51	387	130		116	114	87	105	444	395	141	105	369	682	207	969	130	132
-	1			9			]	-				5	8				1		1			$\begin{bmatrix} & 1 \end{bmatrix}$	1		_	2			1
374	1	2	2	1	104	2	114	312	3	40	13			2	2	3	_	10	1	130	195		_	217	443		442	2	
7500	7501	7502	7503	7504	7505	7506	7507	7508	7509	7510	7511	7512	7513	7514	7515	1216	7517	7518	7519	7520	7521	7522	7523	7524	7525	7526	7527	7528	7529
							emb CAB6298 9.1																						
							(AL022313) dJ1119A7.5 (novel protein (isoform 2)) [Homo sapiens] >sp CAB62989 CAB62989 DJ1119A7.5 (novel protein (isoform 2)) (fragment). Length = 100																						
HCRMX11R	HCRMY28R	HCRMY29R	HCRMY39R	HCRMZ13R	HCRMZ36R	HCRMZ53R	HCRMZ71R	HCRMZ92R	HCRNA39R	HCRNA44R	HCRNA64R	HCRNA88R	HCRNB36R	HCRNB47R	HCRNB56R	HCRNB61R	HCRNB69R	HCRNB77R	HCRNB85R	HCRNC23R	HCRND21R	HCRND28R	HCRND30R	HCRND45R	HCRNE04R	HCRNEIIR	<b>HCRNE15R</b>	HCRNE17R	HCRNE18R
3223	3224	3225	3226	3227	3228	3229	3230	3231	3232	3233	3234	3235	3236	3237	3238	3239	3240	3241	3242	3243	3244	3245	3246	3247	3248	3249	3250	3251	3252

_	Γ							Γ	Γ	ı			_	Γ	_	r —															$\Box$
pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1									
HCRNE34	HCRNE50	HCRNE60	HCRNF01	HCRNF66	HCRNF90	HCRNG33	HCRNG44	HCRNH02	HCRNH78	HCRNI71	HCRN125	HCRNK13	HCRNK40	HCRNK94	HCRNL17	HCRNL38	HCRNL52	HCRNL55	HCRNL60	HCRNL69	HCRNL86	HCRNM46	HCRNM50	HCRNN08	HCRNN11	HCRNN79	HCRNO40	HCRN041	HCRN049	HCRNP05	HCRNP07
											54																				
											40																_	-			
54	146	279	349	453	108	168	561	402	103	483	406	116	464	493	20	202	198	57	311	85	72	122	188	51	141	57	204	471	546	159	183
-	39	-	194	334	1	7	19	175	2	214	59	3	195	200	9	2	100	1	3	2	1	3	45	1	1	1	1	295	274		69
7530	7531	7532	7533	7534	7535	7536	7537	7538	7539	7540	7541	7542	7543	7544	7545	7546	7547	7548	7549	7550	7551	7552	7553	7554	7555	7556	7557	7558	7559	7560	7561
											gb AAF34807. 1 AF2310																				
									PROVIDENCE OF THE PROPERTY OF		(AF231038) SP555 protein [Drosophila melanogaster] Length = 293							:													
HCRNE34R	HCRNE50R	HCRNE60R	HCRNF01R	HCRNF66R	HCRNF90R	HCRNG33R	HCRNG44R	HCRNH02R	HCRNH78R	HCRNI71R	HCRNJ25R	HCRNK13R	HCRNK40R	HCRNK94R	HCRNL17R	HCRNL38R	HCRNL52R	HCRNL55R	HCRNL60R	HCRNL69R	HCRNL86R	HCRNM46R	HCRNM50R	HCRNN08R	HCRNNIIR	HCRNN79R	HCRNO40R	HCRNO41R	HCRNO49R	HCRNP05R	HCRNP07R
3253	3254	3255	3256	3257	3258	3259	3260	3261	3262	3263	3264	3265	3266	3267	3268	3269	3270	3271	3272	3273	3274	3275	3276	3277	3278	3279	3280	3281	3282	3283	3284

pSport1	pSportl	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1																				
HCRNP22	HCRNP34	HCRNP45	HCRNP65	HCRNP76	HCRNQ59	HCRNR03	HCRNR84	HCRNU20	HCRNV70	HCRNV94	HCRNW29	HCRNW34	HCRNW36	HCRNW40	HCRNX03	HCRNX05	HCRNY53	HCRNY85	HCRNZ22	HCRNZ37	HCROB24	HCROB35	HCROB68	HCROB83	HCROB85	HCROD36	HCROE09	HCROE19	HCROE67	HCROE81
																													08	
																													62	
172	315	130	524	114	95	353	198	225	338	298	263	91	182	304	134	103	409	217	492	186	240	261	358	69	504	217	172	218	522	99
2	1	2	390	1	3	84		-	132	2	87	2	42	56	3	2	224	65	181	1	1	1	119	1	193	23	14	30	1	
7562	7563	7564	7565	7566	7567	7568	7569	7570	7571	7572	7573	7574	7575	7576	7577	7578	7579	7580	7581	7582	7583	7584	7585	7586	7587	7588	7589	7590	7591	7592
																													dbj BAA83013 .1	
																													(AB028984) KIAA1061 protein [Homo sapiens] >sp BAA83013 BAA83013 KIAA1061 protein (fragment). Length = 693	
HCRNP22R	HCRNP34R	HCRNP45R	HCRNP65R	HCRNP76R	HCRNQ59R	HCRNR03R	HCRNR84R	HCRNU20R	HCRNV70R	HCRNV94R	HCRNW29R	HCRNW34R	HCRNW36R	HCRNW40R	HCRNX03R	HCRNX05R	HCRNY53R	HCRNY85R	HCRNZ22R	HCRNZ37R	HCROB24R	<b>HCROB35R</b>	HCROB68R	HCROB83R	HCROB85R	HCROD36R	HCROE09R	HCROE19R	HCROE67R	HCROE81R
3285	3286	3287	3288	3289	3290	3291	3292	3293	3294	3295	3296	3297	3298	3299	3300	3301	3302	3303	3304	3305	3306	3307	3308	3309	3310	3311	3312	3313	3314	3315

pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1
HCROE89	HCROF29	HCROF67	HCROF73	HCROG40	HCROG51	HCROG58	HCROG62	HCROG80	HCROH29	HCROH55	HCROH61	HCROH86	HCROI10	HCROI79	HCROI81	HCROI83	HCROJ21	HCROJ35	HCROJ40	HCROJ88	HCROK12	HCROK28	HCROK29	HCROK32	HCROK33	HCROK42	HCROK47	HCROK70	HCROK84	HCROK95	HCROL14	HCROL47
91	2	33	15	132	14	11	11	408	505	7	88	99	266	168	278	376	95	452	155	246	62	250	267	157	150	151	215	131	102	115	244	66
227   406	12 62	149 463	1 105	43   13	2 394	334 561	116 301	112 40	164 50	1 117	2 8	1 6	75 26	1 16	3 27	203 37	15 5	216 45	72   15	1 24	3 6	29 25	103   20	2 15	1 15	2 15	45 21	3 15	1 1(	2 1	113 2	1 9
7593 2	7594	7595 1	7596	7597	7598	7599 3	7600	7601 1	7602 1	7603	7604	7605	. 9092	2092	8092	7609 2	7610	7611 2	7612	7613	7614	7615	7616 1	7617	7618	7619	7620	7621	7622	7623	7624 1	7625
																										·						
HCROE89R	HCROF29R	HCROF67R	HCROF73R	HCROG40R	HCROG51R	HCROG58R	HCROG62R	HCROG80R	HCROH29R	HCROHSSR	HCROH61R	HCROH86R	HCROI10R	HCROI79R	HCRO181R	HCROI83R	HCR0J21R	HCROJ35R	HCROJ40R	HCROJ88R	HCROK12R	HCROK28R	HCROK29R	HCROK32R	HCROK33R	HCROK42R	HCROK47R	HCROK70R	HCROK84R	HCROK95R	HCROL14R	HCROL47R
3316 HCI	3317 HC	3318 HCI	3319 HC	3320 HCI	3321 HCI	3322 HCI	3323 HCI	3324 HCI	3325 HCI	3326 HCI	3327 HCI	3328 HCI	3329 HC	3330 HC	3331 HC	3332 HC	3333 HC	3334 HC	3335 HC	3336 HC	3337 HCI	3338 HCI	3339 HCI	3340 HC	3341 HCI	3342 HCI	3343 HCI	3344 HCJ	3345 HC	3346 HC	3347 HC	3348 HC

HCROL55 pSport1	HCROL69 pSport1	HCROM07 pSport1	HCROM39 pSport1	HCROM50 pSport1	HCROM53 pSport1	HCROM56 pSport1	HCROM63 pSport1	HCROM80 pSport1	HCROM82 pSport1	HCRON01 pSport1	HCRON04 pSport1	HCRON39 pSport1	HCRON42 pSport1	HCRON65 pSport1	HCRON70 pSport1	HCROO20 pSport1	HCROO46 pSport1	HCROP24 pSport1	HCROP51 pSport1	HCROP55 pSport1	HCROP63 pSport1	HCROP69 pSport1	HCROP88 pSport1	HCROQ04 pSport1	HCROQ13 pSport1	HCROQ79 pSport1	HCROQ92 pSport1	HCROR38 pSport1	HCROR69 pSport1	HCROR76 pSport1	HCROR80 pSport1	HCROS08 pSport1
70	6 372	216	59	39	226	0 311	4 256	4 218	100	88	260	108	147	64	88	0 221	9 447	51	6 58	66		06	1 64	85	122	100	325	99	282	. 89 81	09	69 200
7626 2	7627 76	7628	7629 3	7630	7631 2	7632 90	7633 44	7634 24	7635 2	7636 2	7637 3	7638	7639 1	7640 2	7641 2	7642 60	7643 199	7644	7645 26	7646 1	7647 2	7648	7649 11	7650 2	7651 3	7652 2	7653 164	7654	7655	1656	7657	7658 6
HCROL55R	HCROL69R	HCROM07R	HCROM39R	HCROM50R	HCROM53R	HCROM56R	HCROM63R	HCROM80R	HCROM82R	HCRON01R	HCRON04R	HCRON39R	HCRON42R	HCRON65R	HCRON70R	HCRO020R	HCROO46R	HCROP24R	HCROP51R	HCROP55R	HCROP63R	HCROP69R	HCROP88R	HCROQ04R	HCR0Q13R	HCROQ79R	HCROQ92R	HCROR38R	HCROR69R	HCROR76R	HCROR80R	HCROS08R
HCR	HCR	HCRC	HCRC	HCRC	HCR	HCRC	HCRC	HCR	HCRC	HCR(	HCR(	HCR(	HCR(	HCR(	HCR(	HCR	HCR(	HCR(	HCR(	HCR(	HCR	HCR(	HCR	HCR								
3349	3350	3351	3352	3353	3354	3355	3356	3357	3358	3359	3360	3361	3362	3363	3364	3365	3366	3367	3368	3369	3370	3371	3372	3373	3374	3375	3376	3377	3378	3379	3380	3381

2 pSport1	2 pSport1	4 pSport1	5 pSport1	9 pSport1	3 pSport1	'5 pSport1	4 pSport1	4 pSport1	)4 pSport1	% pSport1	54 pSport1	32 pSport1	39 pSport1	58 pSport1	59 pSport1	6 pSport1	8 pSport1	32 pSport1	s8 pSport1	52 pSport1	2 pSport1	9 pSport1	4 pSport1	15 pSport1	58 pSport1	73 pSport1	75 pSport1	6 pSport1	9 pSport1	9 pSport1	1 pSport1	1 pSport1
HCROS22	HCROS52	HCROT14	HCROT15	HCROT19	HCROT23	HCROT75	HCROT84	HCROT94	HCROV04	HCROV08	HCROV64	HCROV82	HCROW39	HCROW68	HCROW69	HCROX16	HCROX18	HCROX32	HCROX38	HCROX52	HCROX92	HCROZ19	HCROZ34	HCROZ45	HCROZ68	HCROZ73	HCROZ75	HCROZ76	HCRPA09	HCRPA19	HCRPA21	HCRPA61
64	109	66	218	227	245	75	236	63	433	229	375	42	63	99	392	105	75	108	105	100	111	363	232	413	129	06	102	397	463	318	187	59
14	2	1	09	09	06	1	3	7	125	32	1	1	1	1	138	-	1	1	1	2	1	184	2	3	1	1	1	224	233	1	38	3
6592	0992	7661	7662	. 7663	7664	2992	9992	1991	8992	6992	0.197	1671	7672	7673	7674	2191	9/9/	<i>LL9L</i>	8191	6191	0892	1892	7682	7683	7684	2892	9892	1897	8892	6892	0692	7691
												: :																				
HCROS22R	HCROS52R	HCROT14R	HCROT15R	HCROT19R	HCROT23R	HCROT75R	HCROT84R	HCROT94R	HCROV04R	HCROV08R	HCROV64R	HCROV82R	HCROW39R	HCROW68R	HCROW69R	HCROX16R	HCROX18R	HCROX32R	HCROX38R	HCROX52R	HCROX92R	HCROZ19R	HCROZ34R	HCROZ45R	HCROZ68R	HCROZ73R	HCROZ75R	HCROZ76R	HCRPA09R	HCRPA19R	HCRPA21R	HCRPA61R
3382	3383	3384	3385	3386	3387	3388	3389	3390	3391	3392	3393	3394	3395	3396	3397	3398	3399	3400	3401	3402	3403	3404	3405	3406	3407	3408	3409	3410	3411	3412	3413	3414

pSport1	pSport1	pSport1	pSport1	pSport1																												
HCRPA91	HCRPB73	HCRPC14	HCRPC30	HCRPC42	HCRPC55	HCRPC56	HCRPC58	HCRPC65	HCRPC80	HCRPC90	HCRPD57	HCRPD85	HCRPE32	HCRPE74	HCRPF41	HCRPF62	HCRPF90	HCRPF92	HCRPG02	HCRPG03	HCRPG11	HCRPG16	HCRPG28	HCRPG37	HCRPG49	HCRPG93	HCRPH31	HCRPH50	HCRPH58	HCRPH93	HCRPI35	HCRPI58
62	59	63	125	113	338	340	121	458	66	247	82	162	174	276	51	209	423	392	116	152	69	211	229	402	164	85	173	117	315	16	217	102
2	15	1	3	15	51	161	2	279	-	83	2	-	1	25		42	223	246	3	3	-	2	95	1	42	2	15	7	92	2	2	1
7692	7693	7694	7695	9692	1697	8692	6692	7700	7701	7702	7703	7704	7705	2706	7707	7708	7709	7710	7711	7712	7713	7714	7715	7716	7717	7718	7719	7720	7721	7722	7723	7724
	:																											-				
HCRPA91R	HCRPB73R	HCRPC14R	HCRPC30R	HCRPC42R	HCRPC55R	HCRPC56R	HCRPC58R	HCRPC65R	HCRPC80R	HCRPC90R	HCRPD57R	HCRPD85R	HCRPE32R	HCRPE74R	HCRPF41R	HCRPF62R	HCRPF90R	HCRPF92R	HCRPG02R	HCRPG03R	HCRPG11R	HCRPG16R	HCRPG28R	HCRPG37R	HCRPG49R	HCRPG93R	HCRPH31R	HCRPH50RA	HCRPH58RA	HCRPH93R	HCRP135RA	HCRP158RA
3415	3416	3417	3418	3419	3420	3421	3422	3423	3424	3425	3426	3427	3428	3429	3430	3431	3432	3433	3434	3435	3436	3437	3438	3439	3440	3441	3442	3443 I	3444 I	3445		3447

pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1												
HCRPI60 p	HCRPI94 p	HCRPJ68 p	HCRPK17 p	HCRPK70 p	HCRPL10 p	HCRPL29 p	HCRPL35 p	HCRPL63 p	HCRPL79 p	HCRPL80 p	HCRPL85 p	HCRPM51 p	HCRPM52 p	HCRPM85 p	HCRPN29 p	HCRPN38 p	HCRPN49 p	HCRPN73 p	HCRPN86 p	HCRPN88 p	HCRPO31 p	HCRPO32 p			HCRPP20 p	HCRPP73 p	HCRPQ23 p	HCRPQ52 p	HCRPQ72 p	HCRPQ79 p
			I	I								H	Ī	1	I	I	Ī		I	I	I		1				1	[	90 I	
	_																						_						88	
221	06	126	354	157	9/	91	145	282	95	235	83	460	<i>L</i> 9	63	19	. 295	249	7.1	143	95	350	58	146	185	59	54	16	63	305	57
54	7	1	202	2	2	2	99	82	3	59	3	26	2	-	2	2	П	3	3	3	09	2	3	6	3	1	2	1	66	7
7725	7726	7727	7728	7729	7730	7731	7732	7733	7734	7735	7736	7737	7738	7739	7740	7741	7742	7743	7744	7745	7746	7747	7748	7749	7750	7751	7752	7753	7754	7755
•				•																									dbj BAA86559 .1	
																													(AB033071) KIAA1245 protein [Homo sapiens] >sp BAA86559 BAA86559 KIAA1245 protein (fragment). Length = 892	
HCRP160RA	HCRP194RA	HCRPJ68RA	HCRPK17R	HCRPK70R	HCRPL10R	HCRPL29R	HCRPL35R	HCRPL63R	HCRPL79R	HCRPL80R	HCRPL85R	<b>HCRPM51R</b>	HCRPM52R	HCRPM85R	HCRPN29R	HCRPN38R	HCRPN49R	HCRPN73R	HCRPN86R	HCRPN88R	HCRP031R	HCRP032R	HCRP069R	HCRPP07R	HCRPP20R	HCRPP73R	HCRPQ23R	HCRPQ52R	HCRPQ72R	HCRPQ79R
3448	3449	3450	3451	3452	3453	3454	3455	3456	3457	3458	3459	3460	3461	3462	3463	3464	3465	3466	3467	3468	3469	3470	3471	3472	3473	3474	3475	3476	3477	3478

pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1																			
HCRPR23	HCRPR62	HCRPR70	HCRPR91	HCRPR95	HCRPS10	HCRPS24	HCRPS50	HCRPT04	HCRPT34	HCRPT78	HCRPT82	HCRPT85	HCRPU09	HCRPU76	HCRPV27	HCRPV39	HCRPV62	HCRPV86	HCRPV91	HCRPW68	HCRPW72	HCRPX21	HCRPX71	HCRPY01	HCRPY59	HCRPY91	HCRPZ13	HCRPZ39	HCRQB75	HCRQC36	HCRQC38
																			100 100												
107	196	161	160	99	136	29	228	144	383	133	352	57	271	312	74	174	534	172	406	98	52	103	161	429	58	218	163	96	257	294	519
3	2	18	17	1	62	2	19	1	279	2	191	1	59	202	ω.	1	340	2	2	3	2	23	3	223	2	9	2	1	33	28	271
7756	7757	7758	7759	09//	1922	7762	7763	7764	7765	99//	1911	21/88	6922	7770	7771	7772	7773	7774	7775	9///	7777	7778	7779	7780	7781	7782	7783	7784	5877	7786	7787
																			dbj BAA91400 .1												
HCRPR23R	HCRPR62R	HCRPR70R	HCRPR91R	HCRPR95R	HCRPS10R	HCRPS24R	HCRPS50R	HCRPT04R	HCRPT34R	HCRPT78R	HCRPT82R	HCRPT85R	HCRPU09R	HCRPU76R	HCRPV27R	HCRPV39R	HCRPV62R	HCRPV86R	HCRPV91R (AK000857) unnamed protein product [Homo sapiens] Length = 180	HCRPW68R	HCRPW72R	HCRPX21R	HCRPX71R	HCRPY01R	HCRPY59R	HCRPY91R	HCRPZ13R	HCRPZ39R	HCRQB75R	HCRQC36R	HCRQC38R
3479 HC		-				-			3488 HC	3489 HC			-			3495 HC	-	-	3498 HC	3499 HC	3500 HC								3508 HC	-	
34	3480	3481	3482	3483	3484	3485	3486	3487	34	34	3490	3491	3492	3493	3494	34	34	3497	34	34	35	3501	3502	35	3504	35	35	3507	35	35	3510

pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSportl	pSport1	ZAP Express																				
HCRQD29	HCRQD41	HCRQD47	HCRQD62	HCRQD75	HCRQF95	HCRQG25	HCRQG72	HCRQ103	HCRQI32	HCRQI34	HCRQI65	HCRQ191	HCRQJ04	HCRQJ08	HCRQJ19	HCRQJ26	HCRQJ54	HCRQJ70	HCRQK15	HCRQL13	HCRQL65	HCRQM37	HCRQM45	HCRQM58	HCRQM59	HCRQM68	HCRQN36	HCRQN42	HCUDT18
						81														i									
						71	_	_							,			_		6	~	~		8				_	10
303	102	418	122	99	207	395	229	164	153	573	54	583	54	06	137	91	48	374	54	462	458	498	162	118	403	293	323	134	235
118	1	2	3	-	_	213	44	3	-	310	-	425	2	1	3	2	-	123	-	307	267		37	2	173	120	210	3	2
7788	6877	7790	1622	7792	7793	7794	7795	9622	7977	7798	7799	7800	7801	7802	7803	7804	7805	9082	7807	7808	7809	7810	7811	7812	7813	7814	7815	7816	7817
						HCRQG25R unnamed protein product [unidentified] Length = 180 emb CAB6919 5.1																-							
HCRQD29R	HCRQD41R	HCRQD47R	HCRQD62R	HCRQD75R	HCRQF95R	HCRQG25R unnamed protein pr	HCRQG72R	HCRQ103R	HCRQ132R	HCRQI34R	HCRQI65R	HCRQ191R	HCRQJ04R	HCRQJ08R	HCRQJ19R	HCRQJ26R	HCRQJ54R	HCRQJ70R	HCRQK15R	HCRQL13R	HCRQL65R	HCRQM37R	HCRQM45R	HCRQM58R	HCRQM59R	HCRQM68R	HCRQN36R	HCRQN42R	HCUDT18R
3511	3512	3513	3514	3515	3516	3517	3518	3519	3520	3521	3522	3523	3524	3525	3526	3527	3528	3529	3530	3531	3532	3533	3534	3535	3536	3537	3538	3539	3540

pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-	pBluescript SK-		pCMVSport 3.0	pCMVSport 3.0	pCMVSport 3.0	pCMVSport 3.0
HCYBA36	HCYBC11	HCYBD19	HCYBE06	HCYBE34	HCYBF65	HCYBH59	HCYBH81	нсувн89	нсувн93	HCYBK65	l. 🕳 🔝	HDPPE11	НБРРИ63	нрорден	нроғу12
				86								88			
				86								98			
189	123	245	227	370	245	240	437	230	694	232	264	355	57	154	482
-	46	54	75	2	3	145	288	357	275	62	91	2	-	2	294
7818	7819	7820	7821	7822	7823	7824	7825	7826	7827	7828	7829	7830	7831	7832	7833
				gb AAF03602. 1								gb AAF26302. 1 AF1846			
				(AF146568) MIL1 protein [Homo sapiens] >sp AAF03602 AAF03602 MIL1 protein. Length = 386								(AF184617) proprotein convertase aPC6C isoform [Branchiostoma californiense] >sp AAF26302 AAF26302 Proprotein convertase aPC6C isoform. Length = 1323			
HCYBA36R	HCYBC11R	HCYBD19R	HCYBE06R	HCYBE34R	HCYBF65R	нсувн59к	HCYBH81R	HCYBH89R	<b>НСҮВН93</b> R	HCYBK65R	HDLAX76R	HDPPE11R	HDPPU63R	HDQDZ61R	HDQFV12R
3541	3542	3543	3544	3545	3546	3547	3548	3549	3550	3551	3552	3553	3554	3555	3556

pCMVSport 3.0	pCMVSport 2.0	pCMVSport 2.0	Uni-ZAP XR	Uni-ZAP XR	Uni-ZAP XR	Uni-ZAP XR	pBluescript	pSport1	pSport1	pSport1	Uni-ZAP XR	Uni-ZAP XR	HFRBW76 Uni-ZAP XR	pBluescript	Uni-ZAP XR	Uni-ZAP XR		pCMVSport 3.0	pCMVSport 3.0
НБQGH10	96SQLQH	HDTMG03	HE2JZ65	HE2RS12	HE2TA21	HE8AE77	HEOAB66	HEONL43	HEOSS64	HFIXZ28	HFKHA60	HFPJM42	HFRBW76	HFVIF71	HGBBA17	HGLAW93	HHEAA94	HHEBR58	ННЕОА63
09	431	409	409	101	305	309	158	76	399	317	440	29	325	232	308	150	141	133	544
-	240	263	245	<u>س</u>	51	202	3	2	283	72	3	2	170	59	96	28	4	2	260
7834	7835	7836	7837	7838	7839	7840	7841	7842	7843	7844	7845	7846	7847	7848	7849	7850	7851	7852	7853
HDQGH10R	HDTDS96R	HDTMG03R	HE2JZ65R	HE2RS12R	HE2TA21R	HE8AE77R	HEOAB66RA	HEONL43R	HEOSS64R	HFIXZ28R	HFKHA60R	HFPJM42R	HFRBW76R	HFVIF71R	HGBBA17R	HGLAW93R	HHEAA94R	HHEBR58R	ННЕQА63R
3557	3558	3559	3560	3561	3562	3563	3564	3565	3566	2958	3568	3569	3570	3571	3572	3573	3574	3575	3576

. 7883 80 253	HKCAA84	HKCAA84   Uni-ZAP XR
7884 60 287	HKCSA76	pBluescript
7885 282 476	HKCSB18	pBluescript
7886 1 240	HKCSB45	pBluescript
7887 2 289	HKCSB47	pBluescript
7888 11 223	HKCSC92	pBluescript
7889 25 357	HKCSF11	pBluescript
7890 2 478	HKCSH46	pBluescript
7891 3 86	HKCSI81	pBluescript
7892 57 479	HKCSJ63	pBluescript
7893 103 225	HKCSL33	pBluescript
7894 1 390	HKCS021	pBluescript
7895 191 487	HKCSP88	pBluescript
7896 170 358	HKCSP90	pBluescript
151	HKCTB29	pBluescript
7898 47 271	HKCTB80	pBluescript
7899 55 354	HKCTD01	pBluescript
7900 81 500	HKCTD27	pBluescript
7901 159 413	HKLAA31	Lambda ZAP II
7902 302 439	HKLAB37	HKLAB37 Lambda ZAP
7903 245 487	HKLAB56	Lambda ZAP II
214	HKLRA55	pBluescript
	HKLRA63	pBluescript
207	HKLRB06	pBluescript
64	HKLRB21	pBluescript
2	HKLRB75	pBluescript
24	HKLSA15	pBluescript
3	HKLSA23	pBluescript
7911 305 427	HKLSA28	pBluescript

HKLSB04 pBluescript		HKLSB41 pBluescript	HKLSB76 pBluescript	HKLSB93 pBluescript		HKLSC42   pBluescript	HKLSD10 pBluescript	HKLSD26 pBluescript	HKLSD61 pBluescript		HKLSD93 pBluescript	HLDCH57 pCMVSport 3.0	HLQFP01 Lambda ZAP	HLYBW70 pSport1	HMWDE95 Uni-ZAP XR	HNBTH48 pSport1	HNBTM76 pSport1			HNTDI71 pCMVSport 3.0	HOCTA23 pSport1	HOCTA91 pSport1	HOCTB04 pSport1	HOCTB19 pSport1	HOCTB32 pSport1	HOCTB56 pSport1	11007005
262   H		388 H	227 H	503 H	337 H		245 H			318 H		436 HI	104 H	413 HI	316 HN	H H H	259 HI	317 H	453 HI	493 H	387 H(					459 H	77
20	-	110	3		77		96		111	169	68	2	18	249	2	41	122	96	46	44	37	1		70	130	1	-
7912	7913	7914	7915	7916	7917	7918	7919	7920	7921	7922	7923	7924	7925	7926	7927	7928	7929	7930	7931	7932	7933	7934	7935	7936	7937	7938	2020
HKLSB04R	HKLSB05R	HKLSB41R	HKLSB76R	HKLSB93R	HKLSC29R	HKLSC42R	HKLSD10R	HKLSD26R	HKLSD61R	HKLSD79R	HKLSD93R	HLDCH57R	HLQFP01R	HLYBW70R	HMWDE95R	HNBTH48R	HNBTM76R	HNOAT40R	HNTCO26R	HNTDI71R	HOCTA23R	HOCTA91R	HOCTB04R	HOCTB19R	HOCTB32R	HOCTB56R	ПОСТВОСВ
	1-	1	_					匚	匚		$\Box$		ļ .	<u> </u>	1	_	_	_				<u> </u>	<u> </u>				F

100   100	HOCTC25 pSport1	HOCTC38 pSport1	HOCTC55 pSport1	HOCTC61 pSport1	HOCTC73 pSport1	HOCTD31 pSport1	HOCTD35 pSport1	-	HOCTD88 pSport1	HOCTD95 pSport1	HOCTE12 pSport1	HOCTE91 pSport1	HOCTF24 pSport1	HOCTF43 pSport1	HOCTF84 pSport1	HODFV69 Uni-ZAP XR	HOEKT71 Uni-ZAP XR	HOEKU65 Uni-ZAP XR	HOGEE21 pCMVSport	HOHAS78 pCMVSport	HOHEE72 pCMVSport	70 HOSNW54 Uni-ZAP XR
7940 7941 7942 7943 7944 7946 7946 7946 7949 7950 7951 7951 7952 7953 7954 7950 7954 7950 7956 7956 7956 7957 7958 7959 7959 7959 7959	100	361	385	138	366	102	264	83	73	102	397	51	61	278	213	210	184	95	383	481	309	
emb CAB7086	2	2	2	1	190	1	172	3	2	1	242	-	2	21	1	64	47	3	60	83	136	20
83	7940	7941	7942	7943	7944	7945	7946	7947	7948	7949	7950	7951	7952	7953	7954	7955	7956	7957	7958	7959	0962	
(AL137661) hypothetical protein [Homo sapiens] >emb CAB70863.1  (AL137661) hypothetical protein [Homo sapiens] >sp CAB70863 CAB70863					:																	emb CAB7086 3.1
																						HOSNW54R (AL137661) hypothetical protein [Homo sapiens] >emb CAB70863.1  (AL137661) hypothetical protein [Homo sapiens] >sp CAB70863 CAB70863 Hypothetical 73.8 kd protein. Length = 661
	3663	3664	3665	3998	3667	3668	3669	3670	3671	3672	3673	3674	3675	3676	3677	3678	3679	3680	3681	3682	3683	3684

3686	HPCRD42R	HPCRD42R (AF151075) HSPC241 [Homo sapiens] Length = 128 [gb AAF36161.   1 AF1510	xF36161.	7963	139	456	82	83	HPCRD42	Other
3687	HPDOA19R			7964	70	306				pSport1
3688	HPFCN76R			2962	06	467			HPFCN76	Uni-ZAP XR
3689	HPJBZ88R	(AL122042) hypothetical protein [Homo sapiens] emb C >pir T34520 T34520 hypothetical protein DKFZp564J157.1 - human (fragment) >sp CAB59179 CAB59179 Hypothetical 17.9 kd protein (fragment). >emb CAB59179.2   (AL122042) hypothetical protein [Homo sapiens] {SUB 22	emb CAB5917 9.1	7966	30	383	94	94	HPJBZ88	Uni-ZAP XR
3690	HRACX76R			1967	2	136			HRACX76	pCMVSport 3.0
3691	HSIFC66R			8962	82	258			HSIFC66	Uni-ZAP XR
3692	HSOBF88R			6962	212	394			HSOBF88	Uni-ZAP XR
3693	HSODE15R			0262	355	444			HSODE15	Uni-ZAP XR
3694	HSVB017R			7971		363			HSVB017	Uni-ZAP XR
3695	HT4CI88R			7972	-	501			HT4CI88	Uni-ZAP XR
3696	HTGEL09R			7973	135	278			HTGEL09	Uni-ZAP XR
3697	HTXRF56R			7974	2	256			HTXRF56	Uni-ZAP XR
3698	HTYND19RA			7975	3	320			HTYND19	pSport1
3699	HTYSJ08Ra			9262	3	236			HTYS108	pBluescript
3700	HWACX88R			7161	3	191		I	HWACX88	pCMVSport 3.0
3701	HWLMA16R			7978	110	253		I	HWLMA16	pSport1
3702	HWLMA24R			7979	2	190		-	HWLMA24	pSport1
3703	HWLMA58R			7980	242	475		H	HWLMA58	pSport1

HWLMA60 pSport1	HWLMA75 pSport1	HWLMA91 pSport1	HWLMB42 pSport1	HWLMC65 pSport1	HWLMC79 pSport1	HWLMD83 pSport1	HWLME13 pSport1	HWLME59 pSport1	HWLME69   pSport1	HWLME71   pSport1	HWLME84   pSport1	HWLMF91 pSport1	HWLMG12 pSport1	HWLMG15 pSport1	HWLMG30 pSport1	HWLMG39 pSport1	HWLMG54 pSport1	HWLMG56   pSport1	HWLMG57 pSport1	HWLMG63 pSport1	HWLMG84 pSport1	HWLMG95 pSport1	HWLMH11 pSport1	HWLMH24 pSport1	HWLMH50 pSport1	HWLM105 pSport1	HWLMI76 pSport1	HWLMJ70 pSport1	HWLMJ80 pSport1	HWLMK20 pSport1	HWLMK25 pSport1	HWLMK31 pSport1
347	458	469	262	204	348	437	195	227	157	387	84	94	111	153	246	106	235	136	249	66	394	216	344	103	220	19	52	247	246	176	66	278
31 198	32 273	191   161		88 58	7986 181	37 3	7988 85	7989 129	7990 2	991 16	7992 1	7993 11	7994 1	1   5662	7996 1	7997 2	7998 38	7999 2	8000 1	8001 1	8002 2	8003 73	8004 216	8005 35	8006 110	8007 2	8008 2	8009 107	8010 1	11 57	8012 1	8013 3
7981	7982	7983	5/_		52	7987	52	52	52	1662	27	51	52	52	52	52	52	52	)8	)8	)8	)8	8(	80	)8	)8	)8	)8	)8	8011	)8	)8
HWLMA60R	HWLMA75R	HWLMA91R	HWLMB42R	HWLMC65R	HWLMC79R	HWLMD83R	HWLME13R	HWLME59R	HWLME69R	HWLME71R	HWLME84R	HWLMF91R	HWLMG12R	HWLMG15R	HWLMG30R	HWLMG39R	HWLMG54R	HWLMG56R	HWLMG57R	HWLMG63R	HWLMG84R	HWLMG95R	HWLMHIIR	HWLMH24R	HWLMH50R	HWLMI05R	HWLMI76R	HWLMJ70R	HWLMJ80R	HWLMK20R	HWLMK25R	HWLMK31R
3704	3705	3706	3707	3708	3709	3710	3711	3712	3713	3714	3715	3716	3717	3718	3719	3720	3721	3722	3723	3724	3725	3726	3727	3728	3729	3730	3731	3732	3733	3734	3735	3736

HWLMK62 pSport1	HWLMM68 pSport1	HWLMM93 pSport1	HWLMN01 pSport1	HWLMN51 pSport1	HWLMP20 pSport1	HWLMP58 pSport1	HWLMP60 pSport1	HWLMP71 pSport1	HWLMQ01 pSport1	HWLMQ73 pSport1	HWLMR23 pSport1	HWLMR69 pSport1	HWLMS31 pSport1	HWLMT42 pSport1	HWLMT57 pSport1	HWLMT64 pSport1	HWLMU07 pSport1	HWLMU13 pSport1	HWLMU26 pSport1	HWLMU41 pSport1	HWLMV34 pSport1	HWLMV60 pSport1	HWLMV66 pSport1		HWLMW93 pSport1	HWLMX13 pSport1	HWLMX67 pSport1	HWLMY52 pSport1	HWLMZ84 pSport1	HWLND18 pSport1	HWLND58 pSport1	HWLND71 pSport1
187	198	48	112	119	202	190	48	63	283	190	1.0	347	103	103	1111	110	143	123	583	203	94	343	66	344	275	285	349	121	208	150	105	158
~	10	_	2	18	2	2		-	104	38	2	3	2	2	1	3	48	4	377	3	2	137	-	204	3	103	158	2	2	7	1	3
8014	8015	8016	8017	8018	8019	8020	8021	8022	8023	8024	8025	8026	8027	8028	8029	8030	8031	8032	8033	8034	8035	8036	8037	8038	8039	8040	8041	8042	8043	8044	8045	8046
(62R	168R	493R	VOIR	451R	220R	58R	260R	71R	201R	273R	KZ3R	(69R	331R	(42R	157R	[64R]	J07R	JI3R	J26R	J41R	/34R	760R	766R	770R	V93R	KI3R	K67R	YS2R	Z84R	)18R	)58R	771R
HWLMK62R	HWLMM68R	HWLMM93R	HWLMN01R	<b>HWLMN51R</b>	HWLMP20R	HWLMP58R	HWLMP60R	HWLMP71R	HWLMQ01R	HWLMQ73R	HWLMR23R	HWLMR69R	HWLMS31R	HWLMT42R	HWLMT57R	HWLMT64R	HWLMU07R	HWLMU13R	HWLMU26R	HWLMU41R	HWLMV34R	HWLMV60R	HWLMV66R	HWLMV70R	HWLMW93R	HWLMX13R	HWLMX67R	HWLMY52R	HWLMZ84R	HWLND18R	HWLND58R	HWLND71R
3737	3738	3739	3740	3741	3742	3743	3744	3745	3746	3747	3748	3749	3750	3751	3752	3753	3754	3755	3756	3757	3758	3759	3760	3761	3762	3763	3764	3765	3766	3767	3768	3769

HWLNF67 pSport1	HWLNF68 pSport1	HWLNG81 pSport1	HWLNH76 pSport1	HWLN143 pSport1	HWLN193 pSport1	HWLNJ40 pSport1	HWLNK59 pSport1	HWLNL41 pSport1	HWLNL71 pSport1	HWLNO22 pSport1	HWLNP11 pSport1	HWLNP22 pSport1	HWLNP43 pSport1	HWLNP65 pSport1	HWLNR24 pSport1	HWLNR26 pSport1	HWLNR27 pSport1	HWLNR48 pSport1	HWLNR57 pSport1	HWLNR81 pSport1	HWLNR83 pSport1	HWLNR92 pSport1	HWLNS19 pSport1	HWLNS37 pSport1	HWLNT23 pSport1	HWLNT40 pSport1	HWLNT48 pSport1	HWLNW90 pSport1	HWLNW92 pSport1	HWLNX01 pSport1	HWLNX64 pSport1	HWLNY25 pSport1
524	73	54	150	66	111	122	75	382	99	190	51	226	250	477	225	176	20	108	153	195	191	51	109		82	177	94	238	88	249	54	238
357	2	1	1	1	25	24	-	2	1	35	1	2	53	322	1	84	3	25	25	136	3	1	44	2	2	1	2	2	2	20	1	2
8047	8048	8049	8050	8051	8052	8053	8054	8055	8056	8057	8028	8059	0908	8061	8062	8063	8064	8065	9908	8067	8908	6908	8070	8071	8072	8073	8074	8075	9/08	8077	8078	8079
3770 HWLNF67R	3771 HWLNF68R	3772 HWLNG81R	3773 HWLNH76R	3774 HWLNI43R	3775 HWLNI93R	3776 HWLNJ40R	3777 HWLNK59R	3778 HWLNL41R	3779 HWLNL71R	3780 HWLNO22R	3781 HWLNP11R	3782 HWLNP22R	3783 HWLNP43R	3784 HWLNP65R	3785 HWLNR24R	3786 HWLNR26R		3788 HWLNR48R	3789 HWLNR57R	3790 HWLNR81R	3791 HWLNR83R	3792 HWLNR92R	3793   HWLNS19R	3794 HWLNS37R	3795 HWLNT23R		3797 HWLNT48R	3798 HWLNW90R	3799 HWLNW92R			3802 HWLNY25R

pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1						
HWLNY40	HWLNY67	HWLNZ75	HWLOA09	HWLOA83	HWLOB93	HWLOC19	HWLOC65	HWLOE46	HWLOF10	HWLOF46	HWLOF52	HWLOF79	HWLOG15	HWLOG17	HWLOG59	HWLO104	HWLOI17	HWLOI25	HWLOI27	HWLOI67	HWLOJ13	HWLOJ19	HWLOJ28	HWLOJ51	HWLOK12	HWLOK36	HWLOK45	HWLOK73	HWLOK90	HWLOL50	HWLOM71
						83 86																									
100	125	152	293	276	73	326	218	125	253	163	105	123	482	150	252	118	414	145	74	363	167	431	126	167	238	54	309	637	336	435	115
2	3	45	3	109	2	156	3	3	119	17	1	1	246	1	46	2	136	71	6	_	12	210	1	9	2	1	154	392	163	238	2
8080	8081	8082	8083	8084	8085	9808	8087	8088	6808	8090	8091	8092	8093	8094	8095	9608	8097	8608	8099	8100	8101	8102	8103	8104	8105	8106	8107	8108	8109	8110	8111
R	R	3	R	R	R	HWLOC19R unnamed protein product [unidentified] Length = 180 emb CAB6919		3	8	8	8	2	R	R	R	~	2	2	2	2	2	2	2	2	R	R	R	R	R	8	R
HWLNY40R	HWLNY67R	HWLNZ75R	HWLOA09R	HWLOA83R	HWLOB93R	HWLOC19R	HWLOC65R	HWLOE46R	HWLOF10R	HWLOF46R	HWLOF52R	HWLOF79R	HWL0G15R	HWLOG17R	HWLOG59R	HWLOI04R	HWL0117R	HWL0125R	HWL0127R	HWLOI67R	HWLOJ13R	HWLOJ19R	HWL0J28R	<b>HWLOJ51R</b>	HWLOK12R	HWLOK36R	HWLOK45R	HWLOK73R	HWLOK90R	HWLOL50R	HWLOM71R
3803	3804	3805	3806	3807	3808	3809	3810	3811	3812	3813	3814	3815	3816	3817	3818	3819	3820	3821	3822	3823	3824	3825	3826	3827	3828	3829	3830	3831	3832	3833	3834

HWLON66 pSport1	HWLON71 pSport1	HWLOO16 pSport1	HWLOQ52 pSport1	HWLOR01 pSport1	HWLOR15 pSport1	HWLOR65 pSport1	HWLOS10 pSport1	HWLOS39 pSport1	HWLOS70 pSport1	HWLOT17 pSport1	HWLOT29 pSport1	HWLOT54 pSport1		HWLOU73 pSport1	HWLOU85 pSport1	HWLOV47 pSport1	HWLOV75 pSport1	HWLOV76 pSport1	HWLOV78 pSport1	HWLOX29 pSport1	HWLOX45 pSport1	HWLOY59 pSport1	HWLOY73 pSport1	HWLOZ31 pSport1	HWLOZ87 pSport1	HWLQA09 pSport1	HWLQA16 pSport1	HWLQA28 pSport1	HWLQA32 pSport1	HWLQA77 pSport1		HWLQC82 pSport1
69	99	51	2 215	180	0 252	1 497	86	63	63	74	186	99	234	89 8	09	63	9 156	6 129	4 152	102 11	06 0	8 496	317	09	9 268	243	316	181	1 258	0 214		0 314
8112 1	8113 1	8114 1	8115 102	8116	8117 10	8118 171	8119 3	8120	8121 1	8122 3	8123 1	8124	8125 1	8126 18	8127	8128	8129 49	8130 46	8131   54	8132 101	8133 10	8134 338	8135 3	8136 1	8137 29	8138	8139 2	8140 2	8141 61	8142 110	8143 53	8144 150
												:																				
														- 4																		
HWLON66R	HWLON71R	HWL0016R	HWLOQ52R	HWLOR01R	HWLOR15R	HWLOR65R	HWLOS10R	HWLOS39R	HWLOS70R	HWLOT17R	HWLOT29R	HWLOT54R	HWLOU39R	HWLOU73R	HWLOU85R	HWLOV47R	HWLOV75R	HWLOV76R	HWLOV78R	HWLOX29R	HWLOX45R	HWLOY59R	HWLOY73R	HWL0Z31R	HWLOZ87R	HWLQA09R	HWLQA16R	HWLQA28R	HWLQA32R	HWLQA77R	_	HWLQC82R
3835	3836	3837	3838	3839	3840	3841	3842	3843	3844	3845	3846	3847	3848	3849	3850	3851	3852	3853	3854	3855	3856	3857	3858	3859	3860	3861	3862	3863	3864	3865	3866	3867

| pSport1  |
|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|
| HWLQD15  | HWLQD16  | HWLQD30  | HWLQD40  | HWLQD42  | HWLQD43  | HWLQD46  | HWLQD89  | НМТОР92  | HWLQD94  | HWLQE28  | HWLQE47  | HWLQE74  | HWLQE83  | HWLQE91  | HWLQF21  | HWLQF47  | HWLQF64  | HWLQF73  | НWLQH32  | HWLQH58  | НМГОН95  | HWLQI87  | HWLQK59  | нмгоме9  | HWLQM91  | HWLQN26  | HWLQN30  | HWLQN44  | HWLQP15  | HWLQP18  | HWLQP26  | HWLQQ83  |
| 307      | 356      | 550      | 228      | 548      | 449      | 274      | 549      | 66       | 505      | 63       | 243      | 99       | 144      | 86       | 97       | 66       | 102      | 187      | 408      | 368      | 102      | 73       | 338      | 89       | 130      | 247      | 173      | 29       | 219      | 124      | 51       | 334      |
| 2        | 3        | 125      | -        | 141      | 3        | 62       | 52       | I        | 182      | 1        | 139      | 1        | 1        | 3        | 2        | 1        | 1        | 2        | 250      | 153      | -        | 2        | 93       | 12       | 44       | 104      | 3        | 2        | _        | 2        | -        | 122      |
| 8145     | 8146     | 8147     | 8148     | 8149     | 8150     | 8151     | 8152     | 8153     | 8154     | 8155     | 8156     | 8157     | 8158     | 8159     | 8160     | 8161     | 8162     | 8163     | 8164     | 8165     | 8166     | 8167     | 8168     | 8169     | 8170     | 8171     | 8172     | 8173     | 8174     | 8175     | 8176     | 8177     |
|          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |
|          |          |          |          | ,        |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |
|          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          | 2        |          |          |          |          |          |          |          |          |          |          |          |          |          |          |
|          |          |          |          | -        |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |          |
| HWLQD15R | HWLQD16R | HWLQD30R | HWLQD40R | HWLQD42R | HWLQD43R | HWLQD46R | HWLQD89R | HWLQD92R | HWLQD94R | HWLQE28R | HWLQE47R | HWLQE74R | HWLQE83R | HWLQE91R | HWLQF21R | HWLQF47R | HWLQF64R | HWLQF73R | HWLQH32R | HWLQH58R | HWLQH95R | HWLQ187R | HWLQK59R | HWLQM69R | HWLQM91R | HWLQN26R | HWLQN30R | HWLQN44R | HWLQP15R | HWLQP18R | HWLQP26R | HWLQQ83R |
| 3868     | 3869     | 3870     | 3871     | 3872     | 3873     | 3874     | 3875     | 3876     | 3877     | 3878     | 3879     | 3880     | 3881     | 3882     | 3883     | 3884     | 3885     | 3886     | 3887     | 3888     | 3889     | 3890     | 3891     | 3892     | 3893     | 3894     | 3895     | 3896     | 3897     | 3898     | 3899     | 3900     |

HWLQR90 pSport1	HWLQR94 pSport1	HWLQT04 pSport1	HWLQT41 pSport1	HWLQT52 pSport1	HWLQT70 pSport1	HWLQU41 pSport1	HWLQU50 pSport1	HWLQU60 pSport1	HWLQW60 pSport1	HWLQW86 pSport1	HWLQX77 pSport1	HWLQY73 pSport1	HWLRB15 pSport1	HWLRC07 pSport1	HWLRC56 pSport1	HWLRE01 pSport1	HWLRE49 pSport1	HWLRF43 pSport1	HWLRF74 pSport1	HWLRH34 pSport1	HWLRH55 pSport1	HWLRH68 pSport1	HWLRL65 pSport1	HWLRM35 pSport1	HWLRM93 pSport1	HWLRN30 pSport1	HWLRN32 pSport1	HWLRO35 pSport1	HWLRO92 pSport1	HWLRP18 pSport1	HWLRP48 pSport1	HWLRP84 pSport1
374	209	51	93	474	69	75	175	103	87	422	71	105	669	69	87	312	381	355	239	08	138	263	264	115	102	66	66	314	155	113	99	66
8178 3	8179 105	8180 1	8181 1	8182 19	8183 1	8184 1	8185 26	8186 2	8187 1	8188 258	8189 9	8190 1	8191 142	8192 1	8193 1	8194 85	8195 235	8196 164	8197 54	8198 3	8199 1	8200 3	8201	8202 53	8203 1	8204 1	8205 1	8206 111	8207 3	8208 6	8209 1	8210 1
				-																												
										:																						
3901 HWLQR90R	3902 HWLQR94R	3903 HWLQT04R		3905 HWLQT52R	3906 HWLQT70R	3907 HWLQU41R	3908 HWLQU50R	3909 HWLQU60R	3910 HWLQW60R	3911 HWLQW86R	3912 HWLQX77R	3913 HWLQY73R	3914 HWLRB15R	3915   HWLRC07R	3916 HWLRC56R	3917 HWLRE01R	3918 HWLRE49R	3919 HWLRF43R	3920 HWLRF74R	3921 HWLRH34R	3922 HWLRH55R		3924 HWLRL65R	3925 HWLRM35R	3926 HWLRM93R	3927 HWLRN30R	3928 HWLRN32R	3929 HWLRO35R	3930 HWLRO92R		3932 HWLRP48R	3933 HWLRP84R

3934	HWLRQ43R	8211	1	66	H	HWLRQ43	pSport1
3935	HWLRR85R	8212	1	51	H	HWLRR85	pSport1
3936	HWLRS49R	8213	3	98	H	HWLRS49	pSport1
3937	HWLRS85R	8214	14	79	H	HWLRS85	pSport1
3938	HWLRT45R	8215	3	65	Η	HWLRT45	pSport1
3939	HWLRT46R	8216	30	116	H	HWLRT46	pSport1
3940	HWLRT77R	8217	68	187	H	HWLRT77	pSport1
3941	HWLRV63R	8218	3	398	H	HWLRV63	pSport1
3942	HWLUF10R	8219		129	H	HWLUF10	pSport1
3943	HWLUF62R	8220	126	281	H	HWLUF62	pSport1
3944	HWLUG36R	8221	2	52	H	HWLUG36	pSport1
3945	HWLUG53R	8222	109	255	H	HWLUG53	pSport1
3946	HWLUG72R	8223	2	145	H	HWLUG72	pSport1
3947	HWLUH57R	8224	3	89	H	HWLUH57	pSport1
3948	HWLUH72R	8225	2	334	H	HWLUH72	pSport1
3949	HWLUH79R	8226	54	179	H	HWLUH79	pSport1
3950	HWLU144R	8227	39	95	H	HWLUI44	pSport1
3951	HWLUISSR	8228	1	312	H	HWLUI55	pSport1
3952	HWLU183R	8229	24	155	H	HWLUI83	pSport1
3953	HWLUJ04R	8230	20	130	H	HWLUJ04	pSport1
3954	HWLUJ19R	8231	1	390	H	HWLUJ19	pSport1
3955	HWLUJ46R	8232	3	137	H	HWLUJ46	pSport1
3956	HWLUL02R	8233	37	186	H	HWLUL02	pSport1
3957	HWLUL39R	8234	1	114	H	HWLUL39	pSport1
3958	HWLUL44R	8235	1	126	H	HWLUL44	pSport1
3959	HWLUL47R	8236	56	202	H	HWLUL47	pSport1
3960	HWLUL65R	8237	30	134	H	HWLUL65	pSport1
3961	HWLUN02R	8238	2	55	H	HWLUN02	p.Sport1
3962	HWLUN03R	8239	1	150	H H	HWLUN03	pSport1
3963	HWLUN23R	8240	1	216	H	HWLUN23	pSport1
3964	HWLUN46R	8241	3	122	H	HWLUN46	pSport1
3965	HWLUN55R	8242	73	138	H H	HWLUN55	pSport1
3966	HWLUN76R	8243	1	117	H	9/NOTMH	pSport1

HWLUN77 pSport1	HWLUN78 pSport1	HWLUN94 pSport1	HWLU012 pSport1	HWLUP38 pSport1	HWLUP63 pSport1	HWLUQ35 pSport1	HWLUQ51 pSport1	HWLUQ54 pSport1	HWLUQ79 pSport1	HWLUQ87 pSport1	HWLUQ94 pSport1		HWLUT21 pSport1	HWLUT89 pSport1	HWLUT94 pSport1	HWLUU23 pSport1	HWLUU88 pSport1	HWLUV35 pSport1	HWLUV39 pSport1	HWLUV67 pSport1	HWLUX01 pSport1	HWLUX69 pSport1	HWLUX81 pSport1	HWLUX84 pSport1	HWLUZ07 pSport1	HWLVA61 pSport1	HWLVA72 pSport1	HWLVA88 pSport1	HWLVB32 pSport1	HWLVB85 pSport1	HWLVD26 pSport1	HWLVD49 pSport1
31 102	59 196	1 54	1 48	12 92	102 221	2 151	2   154	3 98	1 66	149 289	2 178	33 155	1 57	1 81	43 144	2 133	40 192	68 235	1 54	1 72	10 108	2 103	2 85	1 66	2 127	1 144	251 382	4   111	89 229	3 68	31 309	3 113
8244	8245	8246	8247	8248	8249 1	8250	8251	8252	8253	8254	8255		8257	8258	8259	8260	8261	8262	8263	8264	8265	8266	8267	8568	8269	8270		8272	8273	8274	8275	8276
77R	78R	94R	12R	38R	63R	35R	51R	54R	79R	87R	94R	41R	21R	89R	94R	23R	88R	35R	39R	67R	01R	69R	81R	84R	07R	.61R	72R	.88R	32R	85R	26R	49R
HWLUN77R	HWLUN78R	HWLUN94R	HWLU012R	HWLUP38R	HWLUP63R	HWLUQ35R	HWLUQ51R	HWLUQ54R	HWLUQ79R	HWLUQ87R	HWLUQ94R	HWLUR41R	HWLUT21R	HWLUT89R	HWLUT94R	HWLUU23R	HWLUU88R	HWLUV35R	HWLUV39R	HWLUV67R	HWLUX01R	HWLUX69R	HWLUX81R	HWLUX84R	HWLUZ07R	HWLVA61R	HWLVA72R	HWLVA88R	HWLVB32R	HWLVB85R	HWLVD26R	HWLVD49R
3967	3968	3969	3970	3971	3972	3973	3974	3975	3976	3977	3978	3979	3980	3981	3982	3983	3984	3985	3986	3987	3988	3989	3990	1668	3992	3993	3994	3995	3668	3997	3668	3999

HWLVD67 pSport1	HWLVD74 pSport1	HWLVE21 pSport1	HWLVF10 pSport1	HWLVF28 pSport1	HWLVF34 pSport1	HWLVH04 pSport1	HWLVH16 pSport1	HWLVH17 pSport1	HWLVH67 pSport1	HWLV140 pSport1	HWLVI41 pSport1	HWLV115 pSport1	HWLVJ84 pSport1	HWLVK46 pSport1	HWLVK62 pSport1			_	HWLVL71 pSport1		HWLVM05 pSport1	HWLVM23 pSport1	HWLVM49 · pSport1	HWLVN12 pSport1	HWLVN73 pSport1	HWLVN79 pSport1			HWLVR92 pSport1	HWLVS21 pSport1	HWLVS40 pSport1	HWLVT36 psport1
82	89		325	180	443	158	122		132	169	267	186	190	226		06			100	105	129	57	134	231	101				407	69	407	251
8277 2	8278 3	8279 73	8280 170	8281 25	8282 258	8283 3	8284 3	8285 1	8286 1	8287 2	8288 118	8289 1	8290 2	8291 2	8292 3	8293	8294 3	8295 2	8296 2	8297 1	8298 61	8299 1	8300 3	8301 112	8302 3		8304 126	8305 2	8306 162	8307 1	8308 18	8309 3
8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8
HWLVD67R	HWLVD74R	HWLVE21R	HWLVF10R	HWLVF28R	HWLVF34R	HWLVH04R	HWLVH16R	HWLVH17R	HWLVH67R	HWLVI40R	HWLVI41R	HWLVJ15R	HWLVJ84R	HWLVK46R	HWLVK62R	HWLVK88R	HWLVK91R	HWLVL10R	HWLVL71R	HWLVL81R	HWLVM05R	HWLVM23R	HWLVM49R	HWLVN12R	HWLVN73R	HWLVN79R	HWLVR30R	HWLVR40R	HWLVR92R	HWLVS21R	HWLVS40R	HWLVT36R

pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	ort1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	oort1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	p5port1	pSport1
isd	pSt	lSq	isd	pSp	lSq	bSt	lSq	pSt	ISd	Sd	bSt	lSd	Isd	lSq	l pS <sub>I</sub>	Sd	bSı		Sd	bSi	bSi	bS	Sd	bSį		pS			
HWLVV06	HWLVV31	HWLVV64	HWLVV87	HWLVW22	HWLVW49	HWLVW56	HWLVW89	HWLVX39	HWLVX75	HWLVY14	HWLVY55	HWLVY65	HWLVZ12	HWLWA14	HWLWA82	HWLWA91	HWLWB01	HWLWB05	HWLWB42	HWLWB60	HWLWB71	HWLWB73	HWLWB77	HWLWD32	HWLWD56	HWLWD60	HWLWD66	HWLWE25	HWLWE80
																							93				-		
												_											93						
92	259	242	163	174	293	98	58	104	412	011	55	373	53	361	366	123	211	160	142	105	300	156	374	193	343	317	184	316	102
3	14	3	11	1	123	3	20	س	146	3	2	506	3	146	82	1	50	62	2	_	94	-	99	26	149	156	2	95	-
8310	8311	8312	8313	. 8314	8315	8316	8317	8318	8319	8320	8321	8322	8323	8324	8325	8326	8327	8328	8329	8330	8331	8332	8333	8334	8335	8336	8337	8338	8339
																							dbj BAA91151 .1						
																							HWLWB77R (AK000419) unnamed protein product [Homo sapiens] >gb AAF36534.1  (AF154829) 5"(3")-deoxyribonucleotidase [Homo sapiens] {SUB 50-201} Length = 201						
HWLVV06R	HWLVV31R	HWLVV64R	HWLVV87R	HWLVW22R	HWLVW49R	HWLVW56R	HWLVW89R	HWLVX39R	HWLVX75R	HWLVY14R	HWLVY55R	HWLVY65R	HWLVZ12R	HWLWA14R	HWLWA82R	HWLWA91R	<b>HWLWB01R</b>	HWLWB05R	HWLWB42R	HWLWB60R	HWLWB71R	HWLWB73R	HWLWB77R	HWLWD32R	HWLWD56R	HWLWD60R	HWLWD66R	HWLWE25R	HWLWE80R
4033	4034	4035	4036	4037	4038	4039	4040	4041	4042	4043	4044	4045	4046	4047	4048	4049	4050	4051	4052	4053	4054	4055	4056	4057	4058	4059	4060	4061	4062

Γ								Γ				ſ—																			
pSport1	p.Sport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	p.Sport1	pSport1	pSport1	pSport1	pSport1									
HWLWE81	HWLWG36	HWLWH49	HWLWH93	HWLWI26	HWLWI69	HWLWJ36	HWLW137	HWLWK48	HWLWM95	HWLWN12	HWLWN42	HWLWN48	HWLWO57	HWLW064	HWLW078	HWLWP03	HWLWP08	HWLWP13	HWLWP15	HWLWP50	HWLWP87	HWLWQ05	HWLWQ49	HWLWR11	HWLWR26	HWLWR30	HWLWS17	HWLWS19	HWLWS28	HWLWS43	HWLWS64
-	4	-	1				_	14	1	<u> </u>	1		78 F		1		_					I	F	1	I	_	_				
													73																		
149	72	10,4	69	52	166	348	114	39	88	98	51	256	131	223	150	165	98	136	276	414	301	82	146	175	315	247	51	266	179	97	144
3	1	9	1	2	8	172	1	1	2	3	1	20	3	2	1	9/	c	2	1	226	26	2	3	80	196	2	-	3	3	2	-
8340	8341	8342	8343	8344	8345	8346	8347	8348	8349	8350	8351	8352	8353	8354	8355	8356	8357	8358	8359	8360	8361	8362	8363	8364	8365	8366	8367	8368	8369	8370	8371
													dbj BAA91810 .1																		
													HWLWO57R (AK001650) unnamed protein product [Homo sapiens] Length = 216																		
HWLWE81R	HWLWG36R	HWLWH49R	HWLWH93R	HWLWI26R	HWLWI69R	HWLWJ36R	HWLWJ37R	HWLWK48R	HWLWM95R	HWLWN12R	HWLWN42R	HWLWN48R	HWLWO57R	HWLWO64R	HWLWO78R	HWLWP03R	HWLWP08R	HWLWP13R	HWLWP15R	HWLWP50R	HWLWP87R	HWLWQ05R	HWLWQ49R	<b>HWLWR11R</b>	HWLWR26R	HWLWR30R	HWLWS17R	HWLWS19R	HWLWS28R	HWLWS43R	HWLWS64R
4063	4064	4065	4066	4067	4068	4069	4070	4071	4072	4073	4074	4075	4076	4077	4078	4079	4080	4081	4082	4083	4084	4085	4086	4087	4088	4089	4090	4091	4092	4093	4094

HWLWU16 pSport1	HWLWU27 pSport1	HWLWW46 pSport1	HWLWW78 pSport1	HWLWW79 pSport1	HWLWX07 pSport1	HWLWX66 pSport1	HWLWX68 pSport1	HWLXA13 pSport1	HWLXA23 pSport1	HWLXA45 pSport1	HWLXC34 pSport1	HWLXE61 pSport1	HWLXE79 pSport1	HWLXI51 pSport1	HWLXI76 pSport1	HWLXJ59 pSport1	HWLXJ79 pSport1	HWLXK62 pSport1	HWLXN33 pSport1	HWLXO57 pSport1	HWLXO71   pSport1	HWLXO81 pSport1	HWLXP33 pSport1		HWLXP60 pSport1	HWLXQ33 pSport1	HWLXQ50 pSport1	HWLXQ71 pSport1	HWLXQ81 pSport1	HWLXR27 pSport1	HWLXR49 pSport1	HWLXR74 pSport1
240	253	222	234	65	891	64	51	171	226	145	188	3 477	63	54	322	139		7 529	29	211	101	206	69	196	109	52	52	236	110	226	142	240
8372   121	8373 92	8374 1	8375 1	8376 24	8377 19	8378 2	8379 1	8380 1	8381 2	8382 2	8383 36	8384 223	8385 1	8386 1	8387 . 86	8388 2	8389 135	8390 377	8391 2	8392 2	8393 30	8394 3	8395 1	8396 2	8397 2	8398 2	8399 2	8400 3	8401 3	8402 44	8403 2	8404 10
4095 HWLWU16R	4096 HWLWU27R	4097 HWLWW46R	4098 HWLWW78R	4099 HWLWW79R	4100 HWLWX07R	4101 HWLWX66R	4102 HWLWX68R	4103 HWLXA13R	4104 HWLXA23R	4105 HWLXA45R	4106 HWLXC34R	4107 HWLXE61R	4108 HWLXE79R	4109 HWLXISIR	4110 HWLXI76R	4111 HWLXJ59R	4112 HWLXJ79R	4113 HWLXK62R	4114 HWLXN33R	4115 HWLXO57R	4116 HWLXO71R	4117 HWLX081R	4118 HWLXP33R	4119 HWLXP4SR	4120 HWLXP60R	4121 HWLXQ33R	4122 HWLXQ50R	4123   HWLXQ71R	4124 HWLXQ81R	4125 HWLXR27R	4126 HWLXR49R	4127 HWLXR74R

						·	Γ		<u> </u>			Γ_		_				Γ	ı											
pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1	pSport1									
HWLXT31	HWLXV15	HWLXV27	HWLXW17	HWLXW20	HWMBC46	HWMBD22	HWMBD49	HWMBD71	HWMBE31	HWMBE36	HWMBF87	HWMBG63	HWMBG89	HWMBH14	HWMBI08	HWMBI41	HWMBI51	HWMBK47	HWMBL07	HWMBL29	HWMBL57	HWMBL82	HWMBM40	HWMBM51	HWMBM67	HWMBM83	HWMBM87	HWMBN13	HWMBN35	HWMBN52
									72							_														
-	)	(	2	0	~				09 8		~				2	3	- 0	3	2	2	0	9	8		_		9	3	5	₹
284	150	129	115	130	212	433	19	131	408	221	388	54	142	93	315	313	150	313	206	396	330	496	218	10	221	299	136	163	355	274
138	-	-	2	32	114	32	2	3	-	96	209		2	_	1	176		131	9	196	121	17	87	7	81	141	2	2	185	104
8405	8406	8407	8408	8409	8410	8411	8412	8413	8414	8415	8416	· 8417	8418	8419	8420	8421	8422	8423	8424	8425	8426	8427	8428	8429	8430	8431	8432	8433	8434	8435
									gb AAF04012. 1 AF1697																					
									HWMBE31R (AF169797) adaptor protein APPL [Homo sapiens] >sp AAF04012 AAF04012 Adaptor protein APPL. Length = 709																					
HWLXT31R	HWLXV15R	HWLXV27R	HWLXW17R	HWLXW20R	HWMBC46R	HWMBD22R	HWMBD49R	HWMBD71R	HWMBE31R	HWMBE36R	HWMBF87R	HWMBG63R	HWMBG89R	HWMBH14R	HWMBI08R	HWMBI41R	HWMBI51R	HWMBK47R	HWMBL07R	HWMBL29R	HWMBL57R	HWMBL82R	HWMBM40R	<b>HWMBM51R</b>	HWMBM67R	HWMBM83R	HWMBM87R	HWMBN13R	HWMBN35R	HWMBN52R
4128	4129	4130	4131	4132	4133	4134	4135	4136	4137	4138	4139	4140	4141	4142	4143	4144	4145	4146	4147	4148	4149	4150	4151	4152	4153	4154	4155	4156	4157	4158

HWMBN94R HWMBP01R	8436	29	133	HWMBN94 HWMBP01	pSport1 pSport1
HWMBP39R	8438	81	308	HWMBP39	pSport1
HWMBP60R	8439	3	125	HWMBP60	pSport1
HWMBP67R	8440	m	143	HWMBP67	pSport1
HWMBP84R	8441	2	229	HWMBP84	pSport1
HWMBR18R	8442	2	133	HWMBR18	pSport1
HWMBR40R	8443	2	130	HWMBR40	pSport1
HWMBR50R	8444	-	117	HWMBR50	pSport1
HWMBR64R	8445	-	156	HWMBR64	pSport1
HWMBR68R	8446	-	150	HWMBR68	pSport1
HWMBR75R	8447	2	124	HWMBR75	pSport1
HWMBR77R A	8448	2	103	HWMBR77	pSport1
HWMBR79R	8449	_	192	HWMBR79	pSport1
HWMBS06R	8450	-	150	HWMBS06	pSport1
HWMBS28R	8451	2	184	HWMBS28	pSport1
HWMBS75R	8452	3	131	HWMBS75	pSport1
HWMBS87R	8453	19	180	HWMBS87	pSport1
HWMBT23R	8454	43	294	HWMBT23	pSport1
HWMBT71R	8455	52	165	HWMBT71	pSport1
HWMBU43R	8456	91	297	HWMBU43	pSport1
HWMBU67R	8457	1	123	HWMBU67	pSport1
HWMBV48R	8458	49	339	HWMBV48	pSport1
HWMBW45R	8459	3	140	HWMBW45	pSport1
HWMBW54R	8460	26	214	HWMBW54	pSport1
HWMBX10R	8461	23	92	HWMBX10	pSport1
HWMBX94R	8462	3	65	HWMBX94	pSport1
HWMBY09R	8463	24	122	HWMBY09	pSport1
HWMBY34R	8464	65	172	HWMBY34	pSport1
HWMBY51R	8465	1	138	HWMBY51	pSport1
HWMBY90R	8466	2	19	HWMBY90	pSport1
HWMBZ52R	8467	250	411	HWMBZ52	pSport1

HWMBZ60 pSport1	HWMBZ74 pSport1	HWMBZ84 pSport1	HWMCA93 pSport1	HWMCB01 pSport1	HWMCB93 pSport1	HWMCC11 pSport1	HWMCC55 pSport1	HWMCD17 pSport1	HWMCD64 pSport1	HWMCD66 pSport1	HWMCE21 pSport1	HWMCE24 pSport1	HWMCF24 pSport1	HWMCF45 pSport1	HWMCH02 pSport1	HWMCH47 pSport1	HWMCH76 pSport1	HWMCI03 pSport1	HWMCI05 pSport1	HWMCI07 pSport1	HWMCI13 pSport1	HWMCI14 pSport1	_	HWMCI16 pSport1	HWMCI19 pSport1	HWMCI25 pSport1	HWMCI27 pSport1	HWMCI29 pSport1	HWMCI30 pSport1	HWMCI32 pSport1	HWMCI39 pSport1	HWMCI40 pSport1
74	210	161	238	108	233	375	59	232	36	158	49	615	92	127	220	203	140	254	115	213	103	191	124	124	225	126	224	137	211	217	133	112
m	61	9	41	1	3	169	3	95	1	3	2	394	3	2	11	3	3	3	2	1	2	3	2	29	40	_	15	3	2	17	5	20
8468	8469	8470	8471	8472	8473	8474	8475	8476	8477	8478	8479	8480	8481	8482	8483	8484	8485	8486	8487	8488	8489	8490	8491	8492	8493	8494	8495	8496	8497	8498	8499	8500
Z60R	Z74R	Z84R	A93R	BOIR	B93R	CIIR	CSSR	D17R	D64R	D66R	E21R	E24R	F24R	F45R	HO2R	H47R	H76R	103R	105R	107R	113R	114R	115R	116R		125R	127R	129R	130R	132R	139R	TANR
HWMBZ60R	HWMBZ74R	HWMBZ84R	HWMCA93R	HWMCB01R	HWMCB93R	HWMCC11R	HWMCC55R	HWMCD17R	HWMCD64R	HWMCD66R	HWMCE21R	HWMCE24R	HWMCF24R	HWMCF45R	HWMCH02R	HWMCH47R	HWMCH76R	HWMCI03R	HWMCI05R	HWMCI07R	HWMCI13R	HWMCI14R	HWMCI15R	HWMCI16R	HWMCI19R	HWMCI25R	HWMCI27R	HWMCI29R	HWMCI30R	HWMCI32R	HWMCI39R	HWMC140R
4191	4192	4193	4194	4195	4196	4197	4198	4199	4200	4201	4202	4203	4204	4205	4206	4207	4208	4209	4210	4211	4212	4213	4214	4215	4216	4217	4218	4219	4220	4221	4222	4223

HWMCI41 pSport1	HWMCI42 pSport1	HWMCI43 pSport1	HWMCI44 pSport1	HWMCI50 pSport1	HWMCI53 pSport1	HWMCIS5 pSport1	HWMCIS6 pSport1	HWMCI62 pSport1	HWMCI80 pSport1	HWMCI85 pSport1	HWMCI87 pSport1	HWMCI88 pSport1	HWMCI92 pSport1	HWMCJ42 pSport1	· .	HWMCK92 pSport1	HWMCL13 pSport1	HWMCL18 pSport1	HWMCL44 pSport1	HWMCL55 pSport1	HWMCL61 pSport1	HWMCL65 pSport1	HWMCL68 pSport1	HWMCL74 pSport1	HWMCM18 pSport1	HWMCM19 pSport1	HWMCM32 pSport1	HWMCM39 pSport1	HWMCM61 pSport1	HWMCM67 pSport1	HWMCM75 pSport1	HWMCM77 pSport1
199	102	134	143	211	145	169	114	253	102	202	193	001	305	120	204	242	130	145	300	317	162	86	106	162	194	185	179	132	114	104	141	134
2	-	9	9	2	41	2	22	2	-	110	7	2	78	1	13	m	2	7	-	150	43	3	7	_	m	45	72	52	1	3	_	ω
8501	8502	8503	8504	8505	8506	8507	8208	8509	8510	8511	8512	8513	8514	8515	8516	8517	8518	8519	8520	8521	8522	8523	8524	8525	8526	8527	8528	8529	8530	8531	8532	8533
HWMC141R	HWMCI42R	HWMCI43R	HWMC144R	HWMCI50R	HWMCI53R	HWMCISSR	HWMCI56R	HWMCI62R	HWMCI80R	HWMCI85R	HWMCI87R	HWMCI88R	HWMCI92R	HWMCJ42R	HWMCK88R	HWMCK92R	HWMCL13R	HWMCL18R	HWMCL44R	HWMCL55R	HWMCL61R	HWMCL65R	HWMCL68R	HWMCL74R	HWMCM18R	HWMCM19R	HWMCM32R	HWMCM39R	HWMCM61R	HWMCM67R	HWMCM75R	HWMCM77R
HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	HWM	-														
4224	4225	4226	4227	4228	4229	4230	4231	4232	4233	4234	4235	4236	4237	4238	4239	4240	4241	4242	4243	4244	4245	4246	4247	4248	4249	4250	4251	4252	4253	4254	4255	4256

4257	HWMCM80R			8534	3	170			HWMCM80	pSport1
4258	HWMCM85R			8535	2	100			HWMCM85	pSport1
4259	HWMCM89R			8536	3	104			HWMCM89	pSport1
4260	HWMCM92R			8537	8	52			Н ММСМ92	pSport1
4261	HWTBE01R			8538	2	244			HWTBE01	HWTBE01 Uni-ZAP XR
4262	нсорровк			8539	111	_			нсорров	HCQDD08 Lambda ZAP
4263	H2CBK69R	unnamed protein product [Homo sapiens] >emb CAA88750.1  TX protease precursor [Homo sapiens] >gb AAA75171.1  cysteine protease [Homo sapiens] >gb AAA86890.1  Ich-2 [Homo sapiens] >gb AAC99850.1  Mih1/TX isoform alpha [Homo sapiens] >pir A57511 A57511 inte	emb CAA0315 4.1	8540	257	535	86	86	H2CBK69	pBluescript SK-
4264	H2CBD14R	unnamed protein product [unidentified]  >emb CAB41416.1  (AJ238246) sarcolectin [Homo sapiens] >sp Q9Y3R7 Q9Y3R7 SARCOLECTIN.  >emb CAA03727.1  unnamed protein product [unidentified] {SUB 1-135} Length = 469	emb CAA0372 6.1	8541	180	539	88	68	H2CBD14	pBluescript SK-
4265	HCQCJ66R			8542	20	235			НСQСЛ66	HCQCJ66 Lambda ZAP
4266	HCYB053R	HCYBO53R IDN4-GGTR14 PROTEIN. >dbj BAA77334.1  (AB019493) IDN4-GGTR9 [Homo sapiens] {SUB 57-414} >emb CAA22908.1  (AL035303) hypothetical protein [Homo sapiens] {SUB 159-414} Length = 414	sp Q9Y6Y5 Q9 Y6Y5	8543	E.	107	100	100	HCYBO53	pBluescript SK-
4267	HWMCK51R	HWMCK51R unnamed protein product [unidentified] Length = 396 emb CAA0339 6.1	emb CAA0339 6.1	8544	24	284	95	95	HWMCK51	pSport1

Lambda ZAP II	Lambda <b>ZAP</b> II	pCMVS <b>port</b> 2.0	pSport1	pSport1	Uni-ZAP XR	pBluesc <b>ript</b> SK-	pBluescript SK-	Lambda <b>ZAP</b> II	pSport1
<b>НС</b> ССА92	нсорк77	HDTEO77	HCRNC15	HWLRD05	HPWBS43	H2CBU94	H2LAT50	нсосозв	HWMCC56
63		96	66	73	78	89	100	44	
57		87	86	62	92	52	100	32	
231	432	230	539	496	130	360	537	158	n
73	157	72	m	2	2	-	73	30	206
8545	8546	8547	8548	8549	8550	8551	8552	8553	8554
emb CAA2402 7.1		dbj BAA85273	gb AAF37319. 1 AF1021	gb AAD42869. 1 AF1551	dbj BAA03520	emb CAA2056 4.1	gb AAD46135. 1	gb AAA36817. 1	
HCQCA92R URF 2 (NADH dehydrogenase subunit) [Homo sapiens] >gb AAC25441.1  (AF014882) NADH dehydrogenase subunit 2 [Homo sapiens] >gb AAC25443.1  (AF014884) NADH dehydrogenase subunit 2 [Homo sapiens] >gb AAC25444.1  (AF014885) NADH dehydrogenase subunit 2 [Homo sapiens]		NADH dehydrogenase subunit 3 [Pan troglodytes] >dbj BAA85273.1  NADH dehydrogenase subunit 3 [Pan troglodytes] >sp BAA85273 BAA85273 NADH dehydrogenase subunit 3. Length = 115	S	(AF155103) NY-REN-25 antigen [Homo sapiens] >sp[Q9Y5A3 Q9Y5A3 NY-REN-25 ANTIGEN (FRAGMENT). Length = 285	Whole ORF continues from bp19 (right after "tag") to dbj BAA03520 bp1596 ("tga"); similar to chinese hamster 1  sphosphatidylserine synthase. [Homo sapiens] >sp P48651 PSS1_HUMAN PHOSPHATIDYLSERINE SYNTHASE I (SERINE-EXCHANGE ENZYME I) (EC 2.7.8) (KIAA0024). Length	(AL031393) dJ733D15.1 (Zinc-finger protein) [Homo sapiens] Length = 496	n ZNF232 [Homo 46135 Zinc finger	ens] IGER PROTEIN	
HCQCA92R	HCQDK77R	HDTE077R	HCRNC15R	HWLRD05R	HPWBS43R	H2CBU94R	H2LAT50R	HCQCO58R	HWMCC56R
4268	4269	4270	4271	4272	4273	4274	4275	4276	4277

5

10

15

20

25

30

309

PCT/US00/26524

The first column of Table 1 shows the "SEQ ID NO:X" for each of the 4277 polynucleotide sequences of the invention. The second column provides a unique "Sequence/Contig ID" for each sequence.

The third column in Table 1, "Gene Name", provides a putative identification of the gene based on the sequence similarity of its translation product to an amino acid sequence found in a publicly accessible gene database, such as GenBank (NCBI). Methods for determining such sequence similarity are described in Example 1, below. The great majority of the cDNA sequences reported in Table 1 are unrelated to any sequences previously described in the literature. The fourth column in Table 1, "Overlap," provides the database accession no. for the database sequence having similarity.

The preferred translated amino acid sequence, is identified in column five as "AA SEQ ID NO:Y," although other reading frames can also be easily translated using known molecular biology techniques. The polypeptides produced by these alternative open reading frames are specifically contemplated by the present invention. Polynucleotides encoding an amino acid sequence comprising these regions are also embodied, as are polynucleotides which hybridize to polynucleotides encoding these regions.

The sixth and seventh columns in Table 1 provide the location (nucleotide position nos.), "Start" and "End," in the polynucleotide sequence "SEQ ID NO:X" that aligns with homologous database sequence. In one embodiment, the invention provides a polypeptide comprising an amino acid sequence encoded by the portion of SEQ ID NO:X delineated by "Start" and "End". Also provided are polynucleotides encoding such polypeptides.

The eighth and ninth columns provide the "%Id" (percent identity) and "% Si" (percent similarity) observed between the aligned sequence segments of the translation product of SEQ ID NO:X and the database sequence. The eleventh and twelfth columns shown in Table 1 provide a unique Clone identifier (Clone ID:Z) and the Cloning vector contained in the cDNA Clone ID, respectively. At least a portion of SEQ ID NO:X was determined by directly sequencing the referenced clone. The reference clone may have more sequence than described in the sequence listing or the clone may have less. In the vast majority of cases, however, the clone is believed to

310

encode a full-length polypeptide. In the case where a clone is not full-length, a full-length cDNA can be obtained by methods described elsewhere herein.

SEQ ID NO:X and the translated SEQ ID NO:Y are sufficiently accurate and otherwise suitable for a variety of uses well known in the art and described further below. For instance, SEQ ID NO:X is useful for designing nucleic acid hybridization probes that will detect nucleic acid sequences contained in SEQ ID NO:X or the cDNA contained in the deposited clone. These probes will also hybridize to nucleic acid molecules in biological samples, thereby enabling immediate applications in chromosome mapping, linkage analysis, tissue identification and/or typing, and a variety of forensic and diagnostic methods of the invention. Similarly, polypeptides identified from SEQ ID NO:Y may be used to generate antibodies which bind specifically to the colon and/or colon cancer related antigen polypeptides encoded by the cDNA clones identified in Table 1.

5

10

15

20

25

30

Nevertheless, DNA sequences generated by sequencing reactions can contain sequencing errors. The errors exist as misidentified nucleotides, or as insertions or deletions of nucleotides in the generated DNA sequence. The erroneously inserted or deleted nucleotides cause frame shifts in the reading frames of the predicted amino acid sequence. In these cases, the predicted amino acid sequence diverges from the actual amino acid sequence, even though the generated DNA sequence may be greater than 99.9% identical to the actual DNA sequence (for example, one base insertion or deletion in an open reading frame of over 1000 bases).

Accordingly, for those applications requiring precision in the nucleotide sequence or the amino acid sequence, the present invention provides not only the generated nucleotide sequence identified as SEQ ID NO:X and the predicted translated amino acid sequence identified as SEQ ID NO:Y, but also a sample of plasmid DNA containing a human cDNA of the invention deposited with the ATCC, as set forth in Table 1. The nucleotide sequence of each deposited clone can readily be determined by sequencing the deposited clone in accordance with known methods. The predicted amino acid sequence can then be verified from such deposits. Moreover, the amino acid sequence of the protein encoded by a particular clone can also be directly determined by peptide sequencing or by expressing the protein in a

311

suitable host cell containing the deposited human cDNA, collecting the protein, and determining its sequence.

The present invention also relates to vectors or plasmids which include such DNA sequences, as well as the use of the DNA sequences. Table 2 shows the material deposited with the ATCC, the Deposit Date and the ATCC Designation Number.

Table 2

5

10

15

ATCC Deposits	Deposit Date	ATCC Designation Number
LP01, LP02, LP03, LP04,	May-20-97	209059, 209060, 209061, 209062, 209063,
LP05, LP06, LP07, LP08,		209064, 209065, 209066, 209067, 209068,
LP09, LP10, LP11,		209069
LP12	Jan-12-98	209579
LP13	Jan-12-98	209578
LP14	Jul-16-98	203067
LP15	Jul-16-98	203068
LP16	Feb-1-99	203609
LP17	Feb-1-99	203610
LP20	Nov-17-98	203485
LP21	Jun-18-99	PTA-252
LP22	Jun-18-99	PTA-253
PA-005 Phage,	Oct-28-99	PTA-881
PA-005 DNA		PTA-882

each is a mixture of cDNA clones derived from a variety of human tissue and cloned in either a plasmid vector or a phage vector, as shown in Table 9. These deposits are referred to as "the deposits" herein. The tissues from which the clones were derived are listed in Table 9, and the vector in which the cDNA is contained is also indicated in Table 9 as well as Table 1. The deposited material includes the cDNA clones which were partially sequenced and listed in Table 1. Thus, the DNA sequence of Table 1 is only a portion of the sequence included in the clone from which the sequence was derived. Thus, a clone which is isolatable from the ATCC Deposits by

10

15

20

25

30

312

PCT/US00/26524

use of a sequence listed in Table 1 may include the entire coding region of a human gene or in other cases such clone may include a substantial portion of the coding region of a human gene. Although the sequence listing lists only a portion of the DNA sequence in a clone included in the ATCC Deposits, it is well within the ability of one skilled in the art to complete the sequence of the DNA included in a clone isolatable from the ATCC Deposits by use of a sequence (or portion thereof) listed in Table 1 by procedures hereinafter further described, and others apparent to those skilled in the art.

The present invention also relates to the genes corresponding to SEQ ID NO:X, SEQ ID NO:Y, or the deposited clone. The corresponding gene can be isolated in accordance with known methods using the sequence information disclosed herein. Such methods include preparing probes or primers from the disclosed sequence and identifying or amplifying the corresponding gene from appropriate sources of genomic material.

Also provided in the present invention are allelic variants, orthologs, and/or species homologs. Procedures known in the art can be used to obtain full-length genes, allelic variants, splice variants, full-length coding portions, orthologs, and/or species homologs of genes corresponding to SEQ ID NO:X, SEQ ID NO:Y, or the deposited clone, using information from the sequences disclosed herein or the libraries deposited with the ATCC. For example, allelic variants and/or species homologs may be isolated and identified by making suitable probes or primers from the sequences provided herein and screening a suitable nucleic acid source for allelic variants and/or the desired homologue.

Table 3 summarizes the expression profile of polynucleotides corresponding to the clones disclosed in Table 1. The first column provides a unique clone identifier, "Clone ID:Z", for a cDNA clone related to each contig sequence disclosed in Table 1. Column 2, "Library Codes" shows the expression profile of tissue and/or cell line libraries which express the polynucleotides of the invention. Each Library Code in column 2 represents a tissue/cell source identifier code corresponding to the Library Code and Library description provided in Table 5. Expression of these polynucleotides was not observed in the other tissues and/or cell libraries tested. One of skill in the art could routinely use this information to identify tissues which show a

5

10

15

20

25

30

313

PCT/US00/26524

predominant expression pattern of the corresponding polynucleotide of the invention or to identify polynucleotides which show predominant and/or specific tissue expression.

Table 4, column 1, provides a nucleotide sequence identifier, "SEQ ID NO:X," that matches a nucleotide SEQ ID NO:X disclosed in Table 1, column 5. Table 4, column 2, provides the chromosomal location, "Cytologic Band or Chromosome," of polynucleotides corresponding to SEQ ID NO:X. Chromosomal location was determined by finding exact matches to EST and cDNA sequences contained in the NCBI (National Center for Biotechnology Information) UniGene database. Given a presumptive chromosomal location, disease locus association was determined by comparison with the Morbid Map, derived from Online Mendelian Inheritance in Man (Online Mendelian Inheritance in Man, OMIM<sup>TM</sup>. McKusick-Nathans Institute for Genetic Medicine, Johns Hopkins University (Baltimore, MD) and National Center for Biotechnology Information, National Library of Medicine (Bethesda, MD) 2000. World Wide Web URL: http://www.ncbi.nlm.nih.gov/omim/). If the putative chromosomal location of the Query overlapped with the chromosomal location of a Morbid Map entry, the OMIM reference identification number of the morbid map entry is provided in Table 4, column 3, labelled "OMIM ID." A key to the OMIM reference identification numbers is provided in Table 6.

Table 5 provides a key to the Library Code disclosed in Table 3. Column 1 provides the Library Code disclosed in Table 3, column 2. Column 2 provides a description of the tissue or cell source from which the corresponding library was derived.

Table 6 provides a key to the OMIM reference identification numbers disclosed in Table 4, column 3. OMIM reference identification numbers (Column 1) were derived from Online Mendelian Inheritance in Man (Online Mendelian Inheritance in Man, OMIM. McKusick-Nathans Institute for Genetic Medicine, Johns Hopkins University (Baltimore, MD) and National Center for Biotechnology Information, National Library of Medicine, (Bethesda, MD) 2000. World Wide Web URL: http://www.ncbi.nlm.nih.gov/omim/). Column 2 provides diseases associated

314

with the cytologic band disclosed in Table 4, column 2, as determined using the Morbid Map database.

315

PCT/US00/26524

Table 3.

Clone ID	Library Codes
NO: Z	Library Codes
HCENL15	H0052 H0083 H0263 H0620 L0740 L0759 L0777
HSKII86	H0031 H0056 H0090 H0159 H0250 H0264 H0268 H0341 H0422 H0423 H0518
	H0521 H0528 H0575 S0032 S0046 S0132 S0134 S0280 S3014 T0041 T0042
HNHDV16	S0053
HE8BQ01	H0013 H0090 H0263 L0438 L0439 L0521 L0655 L0686 L0731 L0748
	L0750 L0752 L0755 L0766 L0769 L0776 S0148 S0360
НВМСТ70	H0040 H0090 H0421 L0740 L0766
HNTBM67	H0013 H0031 H0032 H0040 H0046 H0052 H0123 H0163 H0170 H0171 H0178
	H0201 H0266 H0355 H0369 H0373 H0381 H0390 H0411 H0427 H0428 H0435
	H0438 H0486 H0519 H0520 H0539 H0550 H0551 H0555 H0562 H0590 H0602
	H0615 H0623 H0624 H0648 H0659 H0660 H0662 H0667 H0670 H0672 H0682
	H0685 H0686 L0005 L0366 L0370 L0372 L0438 L0439 L0471 L0483 L0518
	L0520 L0521 L0526 L0527 L0564 L0565 L0595 L0596 L0598 L0602 L0637
	L0641 L0646 L0650 L0659 L0662 L0663 L0664 L0665 L0666 L0731 L0740 L0751 L0753 L0754 L0755 L0756 L0758 L0759 L0768 L0769 L0771 L0773
	L0731 L0733 L0734 L0733 L0736 L0736 L0736 L0739 L0706 L0709 L0771 L0773
	S0045 S0046 S0049 S0051 S0194 S0196 S0212 S0222 S0242 S0260 S0280
	\$0328 \$0330 \$0354 \$0356 \$0360 \$0370 \$0374 \$0376 \$0380 \$0388 \$0418
}	S0450 S6028 T0006 T0040 T0110
HDPKC15	H0521 S0134 S0300 S0360
HE2OC31	H0170 H0412 H0641 L0759 L0766 L0770 L0775 L0779 S0360
HLWAY38	H0522 H0543 H0553 H0581 L0731 L0740 L0755 L0766 L0771 L0774 L0777
	L0792 L0800 L0803
HBMXT67	H0012 H0052 H0135 H0144 H0171 H0351 H0369 H0457 H0543 H0620
	H0644 H0653 H0658 H0663 L0167 L0438 L0439 L0471 L0526 L0541 L0591
	L0599 L0638 L0646 L0666 L0743 L0747 L0748 L0750 L0754 L0756 L0758
	L0761 L0763 L0764 L0765 L0766 L0770 L0774 L0777 L0779 L0803 L0809
	S0006 S0007 S0010 S0116 S0134 S0360
HCRND41	H0156 H0545 H0587 H0672 L0055 L0663 L0743 L0747 L0752 L0756 L0759
	L0768 L0774 L0775 L0776 L0777 L0783 L0784 S0050 S0278 S0356 S0360
HWLQA43	T0041 H0031 S0150 S0358 S0360
HWLQI33	H0013 H0135 H0163 H0271 H0423 H0549 H0648 L0731 L0740 L0751 L0759
IIWLQ133	L0761 L0764 L0766 L0769 L0770 L0776 L0777 L0779 L0783 L0789 L0796
	L0805 L0806 L0809 S0114 S0126 S0190 S0360 S6024
HSXDD55	L0438 L0439 L0608 L0758 S0036 S0356
HDOPP57	H0522 L0748
HCPAC07	H0340 H0590 H0596 H0641 L0520 L0639 L0745 L0809
HCRNF04	H0171 H0620 H0624 L0592 L0751 L0769 L0774 L0777 S0222 S0356
HMWHN4	H0144 H0341 L0471 L0752 L0766 L0779 S0126 T0110
3	
HTTEL19	H0009 H0031 H0038 H0040 H0041 H0046 H0052 H0059 H0122 H0124 H0144
	H0156 H0250 H0253 H0254 H0255 H0264 H0268 H0392 H0411 H0436
1	H0445 H0478 H0506 H0521 H0543 H0547 H0556 H0563 H0575 H0594
	H0596 H0616 H0620 H0622 H0627 H0650 H0651 H0652 H0657 H0666 L0055
	L0351 L0372 L0382 L0438 L0439 L0456 L0471 L0526 L0543 L0593 L0599
1	L0638 L0646 L0653 L0655 L0659 L0662 L0664 L0665 L0666 L0731 L0740 L0743 L0744 L0747 L0748 L0751 L0754 L0755 L0756 L0757 L0758 L0766
	L0743 L0744 L0747 L0748 L0731 L0734 L0733 L0736 L0737 L0738 L0766 L0769 L0770 L0771 L0775 L0776 L0788 L0794 L0803 L0805 S0026 S0027
	S0038 S0049 S0126 S0132 S0134 S0212 S0222 S0250 S0276 S0278 S0280
	\$0360 \$0376 \$0380 \$0422 \$0424 \$0436 \$0468 \$6028 \$T0002 \$T0006 \$T0042
L	1 2222 2223 20000 20122 20121 20120 20100 20000 10000 10012

	T0067 T0110
HMCFS02	H0170 H0255 H0294 H0423 H0478 H0529 H0539 H0583 H0618 H0656
111.101.502	H0665 H0688 H0702 L0055 L0438 L0483 L0599 L0629 L0636 L0643 L0645
	L0653 L0659 L0665 L0666 L0731 L0749 L0750 L0751 L0754 L0755 L0757
	L0758 L0761 L0764 L0766 L0776 L0779 L0788 L0789 L0790 L0791 L0794
	L0803 L0804 L0805 L0806 L0809 S0282 S0330 S0344 S0420 S0428
HDTBY31	H0004 H0014 H0015 H0032 H0039 H0040 H0052 H0156 H0251 H0266
	H0268 H0318 H0328 H0356 H0361 H0369 H0373 H0375 H0413 H0427
	H0428 H0445 H0486 H0488 H0506 H0519 H0520 H0546 H0551 H0553
	H0555 H0575 H0586 H0587 H0590 H0591 H0594 H0597 H0598 H0601
	H0615 H0622 H0623 H0624 H0631 H0642 H0643 H0644 H0651 H0662
	H0665 H0667 L0163 L0438 L0439 L0471 L0517 L0519 L0527 L0565 L0581
	L0598 L0638 L0654 L0659 L0731 L0740 L0745 L0747 L0748 L0749 L0751
	L0754 L0757 L0758 L0769 L0773 L0776 L0777 L0779 L0804 S0003 S0004
	S0013 S0027 S0028 S0031 S0037 S0040 S0045 S0046 S0126 S0146 S0174
	S0192 S0196 S0208 S0210 S0212 S0214 S0250 S0342 S0356 S0360 S0376 S0390
	S0402 S0418 S0438 S3014 T0067
HTXFI40	H0265 H0444 H0595 L0779 S0376
HADFW62	H0052 H0156 H0333 H0427 H0478 H0521 H0556 H0617 H0646 H0670 L0384
	L0439 L0543 L0591 L0646 L0657 L0745 L0747 L0749 L0756 L0757 L0764 L0769
TIADA (DIC	L0776 S0116 S0210
HARMP12	H0592
HDPCN86	H0309 H0521 S0028 S0356
HFIAX76	H0057 H0529 L0055 L0483 L0750 L0756 L0758 L0759 L0766 L0773 L0776 L0779
HA ED CO2	S0192 S0300 S0360 S0378 S0422 S0452
HAFBC92	H0445 L0740 L0751 T0049
HFIZG43	H0208 H0251 H0445 H0486 H0615 L0439 L0740 L0750 S0214 S0242 T0041
HMEBY61	H0267 T0049
HTJNI76	H0263 H0435 H0486 H0488 H0520 H0579 H0662 H0687 L0438 L0527 L0645 L0656
HWLFM26	L0751 L0753 L0766 L0771 L0779 L0783 L0809 S0192 S0300 S0376 H0085 H0232 H0234 H0597 L0372 L0645 L0789 S0354 S0358 S0374 S0378 S0380
HWLFM26	S0408 S0442
HAQBZ89	H0295 S0218
HWLEH32	S0354
HWLEL81	S0010 S0354 S0356 S0358 S0374 S0432 S0442
HTLHR67	H0013 H0037 H0052 H0187 H0251 H0416 H0509 H0518 H0538 H0543 H0549
III LIINU/	H0551 H0617 H0618 L0362 L0643 L0666 L0717 L0720 L0731 L0748 L0752 L0754
	L0755 L0774 L0775 L0777 L0779 L0789 L0804 S0003 S0010 S0049 S0116 S0280
	S0356 S0360 T0067
HTSGO78	H0039 H0040 H0087 H0131 H0194 H0592 S0001
HCBBA51	H0009 H0013 H0023 H0031 H0039 H0040 H0042 H0044 H0046 H0052 H0087
	H0100 H0125 H0134 H0136 H0144 H0150 H0163 H0170 H0171 H0173 H0177
	H0201 H0204 H0231 H0238 H0255 H0294 H0306 H0309 H0341 H0373 H0393
	H0408 H0411 H0412 H0413 H0421 H0422 H0423 H0428 H0441 H0445 H0486
	H0494 H0546 H0576 H0581 H0586 H0595 H0596 H0597 H0598 H0599 H0606
	H0609 H0616 H0617 H0622 H0633 H0634 H0635 H0646 H0648 H0651 H0653
	H0657 H0658 H0659 H0661 H0663 H0664 H0669 H0670 H0672 H0674 H0682
	H0685 H0686 H0690 L0005 L0163 L0373 L0375 L0394 L0500 L0519 L0520 L0521
	L0522 L0526 L0542 L0588 L0598 L0622 L0623 L0637 L0653 L0731 L0747 L0750
	L0751 L0755 L0757 L0758 L0759 L0761 L0762 L0763 L0764 L0767 L0768 L0769
	L0772 L0773 L0774 L0775 L0782 L0783 L0789 L0803 L0808 L0809 S0003 S0007
	S0011 S0026 S0027 S0031 S0032 S0045 S0046 S0048 S0051 S0053 S0116 S0126
	S0132 S0134 S0142 S0144 S0152 S0188 S0194 S0222 S0260 S0278 S0280 S0282
	S0328 S0330 S0344 S0358 S0360 S0366 S0374 S0376 S0378 S0380 S0388 S0394
	S0422 S0428 S6022 S6024 T0002 T0006 T0023 T0039 T0041 T0048 T0049 T0069
	T0109

TD IMCITED	170020 110040 110052 110125 170144 170104 170252 110200 110250 1T0404 170510
HNTCW73	H0038 H0040 H0052 H0125 H0144 H0194 H0252 H0288 H0359 H0494 H0519
	H0547 H0551 H0657 L0483 S0026 S0027 S0028 S0045 S0046 S0152 S0206 S0342
TIT TIC CO.	S0346 T0103
HLYGG06	H0181 H0444 H0445 H0596 H0657 H0670 L0373 L0439 L0499 L0500 L0502 L0504
]	L0505 L0506 L0507 L0508 L0509 L0511 L0540 L0659 L0663 L0740 L0748 L0750
II A DO A 50	L0752 L0754 L0758 L0763 L0764 L0768 L0769 L0777 L0779 L0783
HAPOA59	H0013 H0038 H0040 H0050 H0056 H0057 H0059 H0144 H0169 H0264 H0266 H0318 H0341 H0428 H0509 H0519 H0529 H0539 H0544 H0556 H0560 H0572
-	H0574 H0575 H0591 H0615 H0616 H0619 H0646 H0648 H0649 H0663 L0096
	L0375 L0378 L0438 L0439 L0471 L0520 L0558 L0588 L0589 L0592 L0593 L0595
	L0601 L0637 L0655 L0659 L0664 L0666 L0731 L0740 L0747 L0748 L0749 L0752
	L0753 L0754 L0756 L0757 L0758 L0759 L0764 L0766 L0768 L0769 L0770 L0771
	L0774 L0775 L0776 L0779 L0780 L0783 L0789 L0794 L0803 L0804 L0805 L0809
ļ	S0014 S0036 S0040 S0132 S0144 S0152 S0250 S0328 S0356 S0392 S0418 S0420
	S0422 S6016 S6024 S6028 T0041 T0067 T0109 T0110
HKLRB18	H0002 H0013 H0014 H0036 H0046 H0050 H0144 H0163 H0234 H0251 H0266
	H0411 H0412 H0413 H0427 H0545 H0550 H0551 H0586 H0593 H0599 H0615
	H0672 L0005 L0163 L0366 L0471 L0542 L0591 L0599 L0659 L0731 L0748 L0750
	L0756 L0757 L0758 L0759 L0777 L0783 L0803 S0026 S0027 S0045 S0152 S0192
	S0206 S0212 S0276 S0328 S0356 S0360 S0418 S3014 T0040
HKAJZ24	H0263 H0494 S0354 S0358 T0039
HJPAU37	H0083 H0097 H0253 H0494 H0556 H0560 H0580 H0593 H0657 L0754 L0766 L0777
	S0356
HHGCU20	H0039 H0052 H0087 H0125 H0135 H0144 H0253 H0318 H0333 H0380 H0445
	H0494 H0542 H0556 H0617 H0624 H0657 H0661 L0471 L0520 L0526 L0622 L0623
	L0731 L0747 L0748 L0749 L0758 L0759 L0764 L0766 L0769 L0774 L0779 L0806
	L0809 S0144 S0210 S0222 S0344 S0360 S0420 S3012 S6022 T0008 T0049 T0082
	T0115
HHEDO80	H0014 H0015 H0156 H0263 H0318 H0411 H0412 H0436 H0445 H0455 H0497
	H0521 H0529 H0542 H0543 H0574 H0575 H0581 H0596 H0599 H0657 H0659
	L0005 L0021 L0455 L0517 L0589 L0590 L0591 L0639 L0664 L0731 L0740 L0741
	L0747 L0752 L0755 L0759 L0766 L0769 L0773 L0775 L0776 L0777 L0780 L0794
1101111770	L0809 S0003 S0132 S0342 S0360 S0374 S0378 S6024
HFIHX78	H0031 H0036 H0051 H0250 H0251 H0263 H0393 H0427 H0436 H0486 H0520 H0575 H0580 H0592 H0596 H0598 H0635 H0661 H0662 L0065 L0373 L0439 L0666
	L0731 L0748 L0752 L0774 L0783 S0194 S0358 S0360 T0023 T0067
HTXOJ32	H0013 H0052 H0056 H0087 H0100 H0150 H0212 H0255 H0352 H0369 H0408
HIAOJ32	H0486 H0556 H0595 H0599 H0619 H0652 H0670 L0352 L0369 L0381 L0415 L0438
	L0439 L0518 L0519 L0528 L0530 L0543 L0588 L0591 L0596 L0605 L0629 L0646
	L0659 L0731 L0741 L0747 L0751 L0757 L0758 L0761 L0764 L0768 L0769 L0771
\	L0773 L0774 L0809 S0003 S0031 S0038 S0045 S0106 S0126 S0134 S0222 S0242
İ	S0250 S0314 S0354 S0356 S0360 S0376 T0010 T0041
HE6FT69	H0100 L0601
HFIHN81	H0012 H0046 H0050 H0051 H0052 H0059 H0090 H0098 H0144 H0170 H0264
	H0309 H0328 H0356 H0370 H0412 H0427 H0428 H0459 H0509 H0521 H0546
	H0547 H0562 H0575 H0591 H0596 H0598 H0616 H0624 H0628 H0634 H0648
	H0658 H0659 H0670 H0672 H0684 L0021 L0157 L0362 L0439 L0444 L0485 L0518
	L0523 L0599 L0600 L0646 L0657 L0659 L0662 L0663 L0664 L0665 L0717 L0731
ĺ	L0738 L0745 L0750 L0751 L0752 L0754 L0756 L0758 L0759 L0766 L0768 L0770
	L0774 L0776 L0777 L0779 L0783 S0003 S0026 S0126 S0194 S0212 S0214 S0222
	S0242 S0260 S0328 S0330 S0354 S0356 S0360 S0376 S0378 S0426 S0464 S6028
	T0006 T0067
HWACZ95	H0012 H0144 H0370 H0393 H0485 H0521 H0574 H0581 H0615 H0620 H0635
	L0381 L0591 L0608 L0648 L0743 L0766 L0774 S0356 S0376
HOELH62	H0040 H0069 H0083 H0090 H0100 H0123 H0144 H0187 H0266 H0333 H0341
	H0370 H0402 H0411 H0413 H0441 H0510 H0525 H0530 H0543 H0544 H0545

318

	H0546 H0580 H0634 L0361 L0375 L0588 L0740 L0747 L0748 L0752 L0767 S0026
	S0040 S0045 S0112 S0114 S0116 S0126 S0182 S0196 S0354 S0358 S0374 S0404
	S0462 S3012
HCE3J64	H0052 H0333 L0439 L0636 L0637 L0742 L0759 S0376 S0388
HWHGE39	H0038 H0051 H0136 H0144 H0178 H0222 H0235 H0305 H0341 H0373 H0393
	H0428 H0435 H0441 H0494 H0506 H0519 H0520 H0542 H0543 H0547 H0555
	H0586 H0587 H0624 H0646 H0648 H0650 H0657 H0667 H0670 H0684 L0021
	L0352 L0438 L0439 L0517 L0586 L0592 L0662 L0664 L0666 L0731 L0740 L0744
	L0747 L0748 L0749 L0752 L0755 L0756 L0757 L0758 L0759 L0766 L0768 L0773
	L0776 L0777 L0779 L0784 L0804 L0809 S0002 S0007 S0010 S0036 S0132 S0192
	S0328 S0356 S0358 S0376 S0424 S0460 T0010 T0115
HNGIN84	H0014 H0036 H0039 H0085 H0183 H0204 H0231 H0506 H0509 H0590 H0596
midnio	H0597 H0622 L0021 L0040 L0364 L0365 L0372 L0373 L0374 L0509 L0596 L0599
	L0646 L0659 L0662 L0761 L0764 L0765 L0772 L0775 L0789 S0052 S0354 S0356
	S0358 S0360 S0374 S0376 S0378 S0380 S0408 S0440 S0442 T0008 T0023 T0109
HPJCI42	H0309 H0370 H0550 H0622 H0624 H0632 H0634 L0005 L0662 L0666 L0764 L0769
HPJC142	
THILL OPE	L0775 L0776 L0794 S0036 S0152
HWLOF51	H0013 H0040 H0318 S0376
HLDOK36	H0012 H0013 H0014 H0015 H0024 H0038 H0039 H0045 H0046 H0051 H0056
	H0059 H0063 H0069 H0074 H0083 H0087 H0090 H0123 H0130 H0135 H0156
	H0163 H0166 H0188 H0213 H0222 H0250 H0261 H0264 H0265 H0266 H0271
	H0294 H0295 H0309 H0316 H0333 H0341 H0351 H0370 H0390 H0393 H0411
	H0416 H0421 H0422 H0423 H0424 H0427 H0428 H0431 H0435 H0436 H0441
	H0444 H0445 H0457 H0478 H0484 H0486 H0494 H0497 H0510 H0519 H0520
	H0521 H0522 H0530 H0539 H0543 H0545 H0546 H0547 H0549 H0551 H0556
	H0574 H0575 H0576 H0580 H0581 H0584 H0585 H0587 H0594 H0595 H0597
	H0598 H0613 H0617 H0618 H0624 H0625 H0633 H0634 H0635 H0638 H0646
	H0648 H0649 H0650 H0652 H0659 H0661 H0662 H0665 H0668 H0672 H0684
	H0695 H0702 L0021 L0361 L0363 L0368 L0369 L0439 L0471 L0517 L0588 L0595
	L0596 L0598 L0637 L0638 L0640 L0649 L0651 L0655 L0659 L0662 L0663 L0665
,	L0666 L0667 L0731 L0740 L0743 L0747 L0748 L0749 L0750 L0751 L0752 L0753
	L0754 L0755 L0756 L0757 L0759 L0761 L0762 L0764 L0766 L0768 L0769 L0770
	L0772 L0774 L0775 L0776 L0777 L0779 L0780 L0782 L0783 L0785 L0787 L0789
	L0794 L0796 L0800 L0803 L0805 S0002 S0003 S0007 S0011 S0026 S0028 S0031
	S0032 S0036 S0040 S0045 S0046 S0049 S0050 S0052 S0114 S0116 S0126 S0132
	S0142 S0144 S0150 S0192 S0194 S0206 S0210 S0212 S0214 S0218 S0222 S0276
1	S0278 S0280 S0306 S0320 S0322 S0330 S0344 S0348 S0350 S0354 S0356 S0358
	S0360 S0372 S0374 S0376 S0378 S0380 S0382 S0384 S0388 S0392 S0422 S0424
	S0426 S0432 S0448 S0450 S0460 S0472 S0474 S3012 S3014 S6022 T0002 T0003
	T0006 T0039 T0042 T0048 T0049 T0109
HSDJF12	H0014 H0031 H0036 H0038 H0085 H0204 H0231 H0263 H0383 H0478 H0487
HSDJF12	H0587 H0590 H0593 H0594 H0596 H0597 H0618 H0619 H0658 H0688 L0040
	L0372 L0374 L0601 L0603 L0645 L0653 L0658 L0659 L0662 L0665 L0666 L0743
	L0748 L0758 L0761 L0764 L0768 L0769 L0770 L0771 L0775 L0780 L0789 L0794
TIMIT EEOO	L0800 L0803 L0804 L0809 S0260 S0280 S0354 S0358 S0374 S0408 S0442 S0458
HWLFF02	H0036 H0056 H0085 H0170 H0171 H0231 H0232 H0263 H0506 H0521 H0590
	H0597 L0040 L0627 L0764 L0765 S0354 S0356 S0358 S0374 S0376
HEMFI21	H0013 H0031 H0046 H0050 H0156 H0212 H0266 H0427 H0509 H0551 H0553
	H0574 H0616 L0455 L0462 L0596 L0747 L0759 S0003 S0046 S0174 S0376
HELGG49	H0030 H0051 H0052 H0135 H0455 H0478 H0617 H0648 H0657 L0375 L0591 L0596
	L0657 L0663 L0665 L0666 L0740 L0743 L0747 L0748 L0752 L0754 L0757 L0800
	S0045 S0046 S0222 S0356 S0376 S0426 S0458 T0067
HRABQ68	H0014 H0052 H0266 H0327 H0370 H0422 H0445 H0519 H0543 H0546 H0555
	H0556 H0615 H0618 L0055 L0438 L0439 L0456 L0598 L0622 L0731 L0740 L0744
	L0745 L0747 L0748 L0750 L0752 L0754 L0755 L0758 L0759 L0761 L0764 L0769
	L0770 L0774 L0776 L0779 L0800 L0803 S0045 S0144 S0356 S0358 S6028 T0071
L	Lacric Barris Barris Bacco Bacco Sacris Barris Basco Bassa Barris Basco Basco Bassa Barris Basco Bassa Barris Basco Basco Bassa Barris Basco Bassa Barris Basco Bas

HCRMP14	S0356
HPRAO21	H0004 H0012 H0024 H0031 H0032 H0156 H0169 H0370 H0428 H0435 H0478
	H0520 H0521 H0596 H0597 H0598 H0644 H0648 H0658 H0664 H0672 H0675
	H0682 L0041 L0471 L0483 L0598 L0599 L0605 L0648 L0657 L0659 L0662 L0663
	L0665 L0666 L0717 L0740 L0744 L0747 L0748 L0749 L0750 L0752 L0754 L0755
	L0758 L0763 L0764 L0768 L0770 L0771 L0774 L0775 L0776 L0779 L0783 L0805
	L0806 L0809 S0328 S0330 S0354 S0358 S0360 S0374 S0376 S0378 S0392 S0448
	T0023 T0079
HAIBU93	H0008 H0012 H0024 H0030 H0031 H0038 H0052 H0068 H0087 H0107 H0135
	H0144 H0178 H0181 H0182 H0194 H0231 H0265 H0266 H0271 H0318 H0320 H0327 H0351 H0369 H0392 H0412 H0416 H0422 H0423 H0445 H0486 H0494
	H0327 H0331 H0369 H0392 H0412 H0416 H0422 H0423 H0443 H0486 H0494
	H0518 H0519 H0521 H0539 H0542 H0543 H0533 H0536 H0561 H0597 H0599 H0606 H0616 H0617 H0620 H0634 H0644 H0646 H0658 H0659 H0663
	H0664 H0673 H0677 H0682 H0684 L0021 L0351 L0352 L0369 L0371 L0372 L0438
	L0439 L0455 L0471 L0517 L0596 L0599 L0623 L0640 L0646 L0648 L0655 L0656
	L0659 L0662 L0664 L0666 L0667 L0717 L0731 L0738 L0744 L0745 L0747 L0748
	L0749 L0751 L0755 L0756 L0757 L0758 L0759 L0761 L0764 L0766 L0768 L0769
	L0770 L0774 L0775 L0776 L0777 L0779 L0788 L0794 S0002 S0038 S0046 S0049
	S0132 S0134 S0142 S0152 S0222 S0280 S0344 S0356 S0358 S0360 S0374 S0376
	S0380 S0388 S0404 S0420 S0428 S6028 T0048 T0068
HHFOV83	H0013 H0014 H0024 H0031 H0032 H0038 H0039 H0040 H0041 H0042 H0046
	H0051 H0052 H0056 H0083 H0108 H0130 H0144 H0156 H0170 H0171 H0208
	H0251 H0316 H0328 H0331 H0341 H0370 H0383 H0386 H0411 H0412 H0428
	H0486 H0494 H0519 H0520 H0538 H0539 H0544 H0546 H0547 H0555 H0576
	H0580 H0581 H0587 H0593 H0598 H0615 H0616 H0617 H0623 H0624 H0633
	H0645 H0651 H0654 H0656 H0657 H0659 H0662 H0663 H0665 H0667 H0670
	H0672 H0673 H0682 H0684 H0686 H0689 L0002 L0005 L0055 L0157 L0367 L0372
	L0439 L0471 L0485 L0517 L0518 L0520 L0521 L0591 L0592 L0593 L0594 L0596
	L0597 L0598 L0599 L0622 L0638 L0648 L0650 L0653 L0655 L0659 L0662 L0663 L0664 L0665 L0666 L0731 L0740 L0745 L0747 L0748 L0749 L0750 L0751 L0752
	L0753 L0754 L0755 L0757 L0758 L0759 L0762 L0764 L0766 L0768 L0769 L0770
	L0771 L0773 L0774 L0775 L0776 L0777 L0779 L0783 L0794 L0806 L0809 S0010
	S0027 S0028 S0036 S0040 S0045 S0046 S0049 S0116 S0126 S0140 S0144 S0148
	S0222 S0250 S0276 S0278 S0282 S0328 S0344 S0356 S0358 S0360 S0376 S0380
	S0390 S0420 S0448 S0472 S3012 S3014 T0008 T0010 T0039 T0040 T0048 T0049
	T0060 T0067 T0110
HMTAE04	H0288 H0341 H0518 H0559 H0560 L0595 S0312
HBBBE52	H0014 H0015 H0031 H0032 H0036 H0038 H0039 H0040 H0050 H0051 H0056
	H0090 H0098 H0119 H0136 H0144 H0147 H0156 H0216 H0264 H0313 H0316
	H0318 H0327 H0328 H0333 H0343 H0359 H0373 H0374 H0375 H0383 H0393
	H0411 H0412 H0413 H0421 H0428 H0431 H0441 H0445 H0461 H0478 H0494
	H0506 H0518 H0519 H0520 H0521 H0542 H0553 H0560 H0574 H0575 H0581
	H0583 H0587 H0593 H0595 H0619 H0622 H0623 H0624 H0626 H0627 H0632
	H0638 H0660 H0661 H0662 H0667 H0683 H0686 H0689 L0021 L0142 L0387 L0389 L0438 L0439 L0471 L0485 L0517 L0518 L0519 L0520 L0540 L0564 L0581 L0590
	L0592 L0604 L0622 L0636 L0638 L0653 L0659 L0662 L0663 L0664 L0665 L0689
	L0731 L0740 L0744 L0747 L0748 L0750 L0752 L0754 L0756 L0757 L0758 L0761
	L0764 L0766 L0768 L0769 L0770 L0771 L0773 L0774 L0775 L0776 L0786 L0803
	L0805 L0809 S0003 S0007 S0010 S0022 S0026 S0031 S0037 S0044 S0046 S0049
	S0050 S0051 S0114 S0126 S0144 S0152 S0210 S0212 S0214 S0222 S0260 S0278
	S0330 S0346 S0354 S0356 S0358 S0360 S0374 S0376 S0390 S0420 S0422 S0424
	S0434 S0438 S0442 S0444 S0474 S6024 S6028 T0040 T0042 T0048 T0067 T0110
	T0115
HDPJL40	H0318 H0436 H0488 H0521 H0553 H0624 H0633 H0667 L0435 L0438 L0439 L0806
	S0002 S0003 S0212 S0358 S0376 T0042
HNTEG54	H0423 H0519 S0045 S0276

320

HNFFZ19	H0059 H0090 H0271 H0436 L0589 S0354
HETCM67	H0004 H0014 H0457 H0529 H0580 H0581 H0638 H0650 L0659 L0666 L0766 L0775
IILI CIVIO	L0793 L0794 S0002 S0003 S0112 S0360 S0374
HTLAK94	H0038 H0052 H0178 H0187 H0253 H0263 H0298 H0399 H0483 H0484 H0510
III La Mesa	H0521 H0545 H0553 H0574 H0590 H0596 H0597 H0615 H0617 H0618 H0622
	H0661 L0439 L0596 L0601 L0740 L0741 L0747 L0748 L0754 L0758 S0001 S0330
	S0358 T0110 T0114
HHETS46	H0266 H0288 H0411 H0421 H0435 H0436 H0543 H0575 H0599 H0617 L0588
	L0590 L0597 L0623 L0731 L0748 L0749 L0753 L0755 L0757 L0758 L0762 L0763
	L0770 L0774 L0775 L0777 L0809 S0028 S0040 S0046 S0049 S0152 S0348 S0380
	S0420 T0048 T0049 T0104
HNAAF81	H0031 H0038 H0040 H0050 H0052 H0252 H0341 H0379 H0402 H0506 H0556
:	H0559 H0586 H0590 H0616 H0622 H0665 L0381 L0439 L0455 L0594 L0608 L0717
	L0749 L0758 S0010 S0040 S0126 S0144 S0356 S0358 S0380 T0041 T0110
HRDFP67	H0031 H0124
HDPPM58	H0031 H0046 H0052 H0090 H0124 H0144 H0244 H0266 H0375 H0421 H0494
	H0506 H0520 H0521 H0522 H0575 H0580 H0590 H0591 H0619 H0622 H0644
	H0672 L0439 L0456 L0471 L0563 L0592 L0593 L0595 N0007 N0009 S0002 S0003
	S0007 S0028 S0036 S0040 S0045 S0142 S0150 S0278 S0346 S0356 S6028 T0042
LITATION2	T0110 T0115 H0052 H0063 H0069 H0135 H0156 H0284 H0288 H0333 H0341 H0423 H0445
НТАНС93	H0052 H0063 H0069 H0133 H0136 H0284 H0288 H0333 H0341 H0423 H0443 H0542 H0543 H0556 H0586 H0590 H0592 H0634 H0635 L0596 L0758 L0766 S0046
	S0126 S0150 S0358 S0364
HCRNJ44	H0069 H0271 H0635 S0356
HWLNH36	H0008 H0032 H0056 H0171 H0328 H0352 H0412 H0510 H0624 H0648 H0658
11 W EI VII 30	H0659 L0005 L0142 L0143 L0591 L0593 L0599 L0605 L0662 L0666 L0740 L0742
	L0751 L0754 L0756 L0758 L0774 L0777 S0007 S0356 S0376
HNSMB24	H0483 H0658 L0638 L0657 L0664 L0665 L0762 L0771 L0779 L0783 S0376
HLLCC54	H0052 H0265 H0354 H0423 H0556 H0575 L0756 S0002 S0376
HTTIU53	H0634 S0376
HCYBG26	H0144 H0521 H0644 L0750 T0114
HTODN93	H0264 H0556 H0581 S0356
HWLQH59	L0754 L0755 L0777 S0360
HTTFP72	H0024 H0040 H0169 H0333 H0422 H0486 H0543 L0601 L0662 S0134 S0354 S0420
HWLIN80	H0263 H0309 H0615 L0021 L0040 L0595 L0662 L0666 L0683 L0731 L0740 L0747
	L0748 L0757 L0759 L0761 L0766 L0773 L0774 L0803 L0809 S0356 S0358
HKMLN95	H0098 H0144 H0147 H0169 H0171 H0305 H0328 H0333 H0413 H0423 H0431
	H0445 H0494 H0519 H0521 H0522 H0543 H0544 H0547 H0574 H0586 H0590 H0632 H0643 H0648 H0650 H0657 H0660 H0661 L0363 L0367 L0372 L0438 L0439
	L0471 L0526 L0544 L0545 L0586 L0592 L0593 L0596 L0599 L0605 L0646 L0648
	L0659 L0664 L0731 L0740 L0747 L0748 L0749 L0752 L0753 L0754 L0755 L0756
	L0757 L0758 L0759 L0766 L0775 L0776 L0794 L0803 L0805 S0003 S0036 S0040
	S0051 S0106 S0144 S0212 S0214 S0276 S0330 S0354 S0356 S0358 S0360 S0374
	S0422 S0450 S6028 T0002 T0110
HDPAM86	H0031 H0038 H0050 H0052 H0056 H0170 H0194 H0265 H0266 H0274 H0286
	H0288 H0292 H0316 H0318 H0341 H0375 H0393 H0412 H0435 H0437 H0445
	H0453 H0457 H0486 H0494 H0519 H0521 H0542 H0543 H0545 H0556 H0560
	H0561 H0580 H0587 H0591 H0615 H0644 H0645 H0657 H0658 H0659 H0661
	H0662 L0372 L0389 L0438 L0439 L0455 L0471 L0588 L0591 L0608 L0637 L0647
	L0650 L0657 L0659 L0665 L0666 L0717 L0740 L0743 L0747 L0748 L0749 L0752
	L0754 L0758 L0759 L0761 L0766 L0768 L0769 L0771 L0774 L0776 L0777 L0803
	L0805 S0002 S0010 S0011 S0152 S0218 S0250 S0328 S0354 S0356 S0358 S0360
	S0370 S0376 S0380 S0418 S0420 S0424 S0426 T0039 T0042 T0110   H0038 H0266 H0328 H0520 L0438 L0471 L0752 L0766 S0003 S0192 S0374 S0438
	1 - DODZY DOZAK DOZIY DOSIO LOZIY LOZIZ LOZIZ LOZIKI 1075 LOZIAK SODOZ SODOZ SODI
HNTMD17	
HNTMD17 HHFJL44	H0036 H0206 H0328 H0320 L0438 L0471 L0732 L0700 30003 30192 30374 30438 H0031 H0309 H0355 H0509 H0510 H0593 H0595 H0619 H0624 H0670 L0021 L0366 L0438 L0731 L0744 L0750 L0752 L0754 L0755 L0758 L0759 L0761 L0767

	L0770 L0774 L0776 L0777 L0794 L0803 L0804 L0805 S0026 S0028 S0142 S0148
	S0196 S0276 S0356 S0360
HNSAD12	H0024 H0069 H0144 L0740 S0434
HWMBM1	H0013 H0036 H0038 H0052 H0590 H0596 H0616 L0040 L0596 L0646 L0654 L0659
3	L0755 L0758 L0764 L0766 L0768 L0771 L0775 L0777 L0794 L0803 S0354 S0358
	S0376 S0404 S0458 T0109
HBXCF35	H0494 L0757 L0766 S0038
HWLFG75	H0205 H0539 H0688 L0142 L0747 L0759 L0770 S0049 S0051 S0260 S0278 S0354
	S0356 S0376 S3014
HWAGI58	H0013 H0144 H0170 H0171 H0428 H0445 H0518 H0538 H0542 H0581 H0592
	H0596 H0619 H0620 H0622 H0624 L0002 L0163 L0439 L0485 L0521 L0532 L0608
	L0646 L0666 L0731 L0752 L0754 L0756 L0759 L0766 L0775 L0777 L0786 S0003
	S0214 S0346 S0360 S0374 S0422
HNGGK17	H0011 H0012 H0024 H0036 H0051 H0100 H0131 H0135 H0144 H0253 H0265
	H0266 H0333 H0341 H0352 H0355 H0393 H0437 H0509 H0519 H0521 H0545
	H0546 H0551 H0553 H0561 H0580 H0581 H0586 H0619 H0620 H0631 L0352
	L0369 L0384 L0483 L0512 L0524 L0534 L0596 L0597 L0603 L0659 L0662 L0664
	L0731 L0740 L0744 L0747 L0750 L0751 L0755 L0758 L0761 L0764 L0766 L0769
	L0770 L0774 L0775 L0777 L0779 L0794 S0002 S0028 S0031 S0038 S0040 S0045
	S0052 S0126 S0206 S0276 S0278 S0356 S0358 S0360 S0374 S0376 S0436 S3014
**************************************	T0002 T0110
HCRQG35	L0752 S0356 S0376
HCRQM44	H0615 L0493 L0519 L0526 L0776 L0794 S0356 S0360
H2CBN90	H0144 H0435 H0485 H0633 L0455 L0738 S0032 S0360 T0110
HTTDU01	H0032 H0038 H0040 H0412 H0488 H0520 H0598 H0626 H0664 L0369 L0485 L0593
·	L0626 L0638 L0662 L0666 L0756 L0758 L0759 L0774 L0779 L0791 L0794 L0803
**********	L0804 S0003 S0010 S0026 S0040 S0356 T0002 T0006 T0110
HAHCU44	H0052 H0176 H0599 H0672 L0752 S0051 S0356 S0374 S0378 S0380
HOSPA23	H0036 H0316 H0436 H0478 H0596 H0622 H0657 H0661 H0666 L0439 L0591 L0664
	L0740 L0752 L0754 L0756 L0759 L0766 L0779 L0794 L0803 S0003 S0040 S0049
VIO CDICI	S0150 S0358 S0422 S0444
H2CBI61	H0013 H0038 H0039 H0305 H0479 H0510 H0617 H0622 H0646 H0658 L0352
	L0371 L0388 L0439 L0485 L0591 L0595 L0596 L0655 L0755 L0756 L0758 L0759 L0761 L0764 L0766 L0769 L0779 S0330 S0346 S0364 T0042 T0110
HCOCD 67	
HCQCR67 HCROV23	H0596   S0356
HCRMZ75	H0031 H0318 H0509 H0581 H0628 H0665 L0595 L0596 L0599 L0600 L0608 L0740
HCKWIZ/3	L0754 S0126 S0328 S0356 S0360 T0042 T0049
HCRMZ85	H0090 H0421 H0509 L0055 L0369 L0591 L0748 L0749 L0752 L0759 L0766 L0771
nCRV1203	L0772 L0794 S0116 S0176 S0356 S0420
HCROM08	S0356
HCRNM87	\$0356
HDPGS84	H0521
	H0024 H0100 H0251 H0393 H0445 H0543 H0553 H0644 H0664 L0517 L0588 L0596
HE6BJ48	L0646 L0738 L0740 L0754 S0218 S0276 S0356
HEOOMALI	H0144 H0239 H0435 H0455 H0661 H0663 H0672 L0438 L0591 L0657 L0659 L0731
HE9QM31	L0740 L0752 L0756 L0759 L0775 L0779 S0007 S0356 S0374 T0110
HDTMC78	H0351 H0486 H0648 H0658 L0655 L0665 L0731 L0747 L0752 L0754 L0756 L0758
HDTMC/8	L0774 L0776 L0779 L0805 L0809 S0242 S0356 S0374 S0424 T0041
HOSMQ26	H0068 H0123 H0263 H0351 H0423 H0542 H0659 H0663 L0362 L0372 L0588 L0591
nosmy26	
	L0655 L0747 L0750 L0752 L0753 L0757 L0764 L0766 L0773 L0776 L0779 S0003
HEADYON	S0045 S0276 S0376 S3014 T0002   H0144 H0156 H0486 H0494 L0520 L0731 L0754 L0758 S0360 T0003 T0109
HKABV02	
HKACE03	H0038 H0294 H0494 H0580 H0620 H0685 S0356 S0420
HBIOR20	H0547 H0593 H0597 L0611 L0809

НКАНЈ56	H0031 H0393 H0494 H0519 H0521 H0581 L0601 L0602 L0774 S0028 S0045 S0376 T0042 T0049 T0082
HCVDV40	H0013 H0038 H0039 H0063 H0144 H0169 H0250 H0252 H0271 H0295 H0331
HSYDX40	H0355 H0393 H0486 H0521 H0538 H0542 H0551 H0553 H0575 H0590 H0593
	H0617 H0648 H0676 H0684 H0690 L0054 L0455 L0499 L0545 L0599 L0654 L0655
	L0731 L0740 L0747 L0748 L0750 L0757 L0758 L0759 L0763 L0764 L0766 L0768
	L0769 L0770 L0774 L0775 L0776 L0779 L0809 S0015 S0027 S0036 S0046 S0126
	S0150 S0212 S0260 S0356 S0360 S0378 S0418 T0041
HMTAD91	H0039 H0184 H0518 H0521 H0560 H0650 H0659 H0675 L0655 L0666 L0731 L0758
	L0766 S0142 S0182 S0260 S0330 S0356
HDPTI77	H0178 H0306 H0402 H0412 H0428 H0521 L0366 L0438 L0605 L0659 L0742 L0747
	L0756 L0758 L0761 L0764 L0771 L0777 L0780 L0794 S0222 S0354 S0360
HFKHN59	H0135 H0596 H0620 H0659 L0055 L0157 L0438 L0591 L0746 L0747 L0748 L0750
	L0752 L0754 L0756 L0757 L0758 L0759 L0764 L0766 L0771 L0774 L0777 L0780
	L0804 L0805 L0806 S0144 S0364 S0376
HWLHW1	H0036 H0506 L0194 S0354 S0358 S0360 S0440 T0115
9	
HTWBQ51	H0436 H0657 H0670 L0352 L0438 L0439 L0593 L0598 L0740 L0747 L0752 L0754
	L0758 L0759 L0776 L0777 S0374 S0376
HTFNM11	S0242 S0424
HCQBI18	H0014 H0059 H0204 H0252 H0263 H0617 H0643 H0682 H0684 L0362 L0374 L0382
	L0642 L0659 L0662 L0665 L0748 L0749 L0764 L0769 L0770 L0771 L0772 L0775
	L0809 S0328 S0358
HCRPJ86	H0351 H0648 H0657 H0659 H0661 L0439 L0651 L0665 L0747 L0752 L0766 L0777
	L0779 S0354 S0356 S0358
HKLAA30	H0184 H0393 H0509 L0471 L0777
HIBEL82	L0439 L0731 L0748 S0356 T0010
HBIBQ89	H0052 H0266 H0318 H0333 H0392 H0402 H0521 H0522 H0556 H0581 H0620
1	H0662 L0021 L0351 L0438 L0439 L0471 L0588 L0659 L0731 L0751 L0756 L0758
•	L0766 L0770 L0772 L0776 L0779 L0792 S0011 S0027 S0049 S0222 S0282 S0418 T0008
HDPIG12	H0380 H0521 S0356
HTNBJ90	H0013 S0376 T0067
HCRON75	S0356
HDQEG93	H0013 H0370 H0494 H0521 H0689 L0596 L0659 S0354 S0356
HPLBS64	H0030 H0184 H0428 H0574 L0351 L0438 L0439 L0596 L0655 L0747 L0748 L0757
HI LD304	L0761 L0766 L0779 S0122 S0132 S0212 S0356 T0008
HCQDT68	H0506 H0596 L0748 S0026 S0356
HWBCW8	H0251 H0266 H0486 H0510 H0539 H0574 H0575 H0580 H0632 L0599 L0740 L0744
0	L0748 L0766 L0774 L0775 L0777 L0779 L0805 S0028 S0356 S0360
H2CBI34	H0031 H0266 H0542 H0574 L0581 L0759 L0806 S0031 S0150 S0278 T0110
HILCJ69	H0031 H0090 H0445 H0555 H0581 H0669 L0438 L0520 L0598 L0740 L0766 S0152
	S0212 S0330 S0376 S0422 T0002
HUSGQ41	H0412 S0222 S0356
HCROG59	H0024 H0050 H0068 H0123 H0150 H0251 H0328 H0546 H0549 H0644 H0658
	H0689 L0603 L0648 L0659 L0717 L0731 L0746 L0747 L0749 L0750 L0758 L0764
	L0774 L0777 L0779 L0794 L0803 L0805 L0809 S0206 S0280 S0356 T0082
HBGNK79	H0131 H0402 H0617 L0751 L0794
HCRNH72	H0013 H0171 L0593 S0356
HDTBL01	H0039 H0081 H0086 H0087 H0136 H0144 H0156 H0170 H0231 H0263 H0295
	H0318 H0327 H0331 H0341 H0370 H0375 H0393 H0413 H0421 H0445 H0483
	H0486 H0494 H0521 H0522 H0542 H0544 H0545 H0547 H0574 H0580 H0581
	H0586 H0591 H0596 H0599 H0615 H0625 H0631 H0650 H0660 H0661 H0662
J	H0670 H0690 L0163 L0384 L0521 L0527 L0588 L0608 L0637 L0659 L0731 L0740
	L0742 L0747 L0748 L0749 L0750 L0751 L0752 L0754 L0755 L0757 L0758 L0759

	107(1) 107(4) 107(7) 107(2) 107(2) 107(1) 107(1) 107(1) 107(1)
	L0761 L0764 L0766 L0768 L0769 L0770 L0771 L0773 L0774 L0775 L0776 L0786
	L0794 L0809 S0028 S0036 S0045 S0342 S0358 S0360 S0366 S0374 S0376 S0380
	T0042 T0049 T0082
H2CBP05	H0046 H0327 H0594 L0193 L0748 L0764 L0766 T0110
HTJMN69	H0235 H0411 H0488 H0547 L0771
HCQDM23	H0085 H0204 H0231 H0232 H0235 H0263 H0489 H0596 H0597 H0688 L0646 L0764 L0771 S0354 S0358 S0376 S0406
HTFNZ86	H0014 H0040 H0266 H0316 H0486 H0519 H0542 H0551 H0556 H0561 H0580
HIFNZ80	H0590 H0616 H0648 H0682 L0163 L0564 L0657 L0659 L0748 L0754 L0758 L0766
	L0777 S0116 S0356 S0360 S0376 S0424
HUSXP66	H0413 H0543 H0657 L0626 L0747 L0748 L0749 L0759 L0766 L0780 S0356
HOFMV44	H0415 H0539 H0598 L0518 L0740 L0749 L0766 L0774 L0777 L0779 S0356
HSLJN60	H0031 H0263 L0599 L0608 L0644 L0666 L0731 L0779 S0250 S0390
	H0031 H0203 L0399 L0008 L0044 L0000 L0731 L0779 30230 30390 H0542 L0596 L0770 L0779 T0110
HHECM62	H0497 H0521 H0563 H0615 L0746 L0752 L0763 L0774 L0803 S0376 S6028 T0041
HMIAQ09	T0067
HMUBG89	H0266 H0529 H0672 L0439 L0740 L0748 L0749 L0750 L0776 L0794 L0803 L0805
111102005	S0360
HCRQC94	H0004 H0014 H0032 H0038 H0039 H0040 H0052 H0059 H0090 H0144 H0187
	H0242 H0331 H0421 H0422 H0435 H0486 H0492 H0510 H0519 H0521 H0522
	H0538 H0539 H0542 H0543 H0553 H0581 H0591 H0593 H0622 H0646 H0654
	H0658 H0659 H0664 H0666 H0684 H0686 L0021 L0105 L0438 L0439 L0471 L0485
	L0560 L0590 L0591 L0600 L0637 L0652 L0659 L0664 L0665 L0666 L0667 L0731
	L0740 L0743 L0745 L0746 L0747 L0748 L0749 L0750 L0754 L0757 L0758 L0759
	L0761 L0764 L0766 L0768 L0771 L0774 L0775 L0776 L0777 L0779 L0784 L0794
	L0803 L0804 L0805 L0806 L0809 S0003 S0010 S0122 S0142 S0152 S0192 S0194
	S0222 S0242 S0330 S0356 S0358 S0360 S0374 S0376 S0414 S0426 S6024 T0082
HUCPH16	H0040 H0052 H0329 H0581 H0586 L0646 L0740 L0748 L0749 L0758 L0779 S0003
	S0013 S0114 S0136 S0356 S0376 S0420 S0426
HCRMN10	H0135 H0331 S0049 S0356
HCGBB63	H0009 H0012 H0013 H0052 H0063 H0081 H0083 H0090 H0144 H0170 H0263
	H0318 H0373 H0393 H0445 H0458 H0494 H0543 H0547 H0549 H0556 H0583
	H0596 H0623 H0624 H0638 H0657 H0659 H0662 H0668 H0670 H0672 L0021
	L0163 L0375 L0381 L0382 L0386 L0471 L0520 L0651 L0664 L0742 L0747 L0748 L0749 L0750 L0752 L0755 L0756 L0757 L0758 L0759 L0763 L0764 L0767 L0768
	L0749 L0730 L0732 L0733 L0736 L0737 L0738 L0739 L0703 L0704 L0707 L0708 L0769 L0771 L0773 L0774 L0775 L0776 L0783 S0003 S0007 S0027 S0032 S0046
	S0276 S0330 S0358 S0360 S0378 S3014 T0006 T0040 T0041
HWLLU74	H0013 H0030 H0031 H0032 H0135 H0411 H0590 H0597 H0682 L0438 L0439 L0666
nwllo/4	L0731 L0743 L0744 L0747 L0750 L0758 L0759 L0763 L0764 L0769 L0772 L0773
	L0776 L0777 S0046 S0053 S0112 S0222 S0350 S0356 S0358 S0360 S0376 S0378
HCUEB32	H0402 H0620 S0360
HCRNA26	H0445 H0478 H0658 L0741 L0803 S0356
HCRMG60	H0059 H0575 L0759 S0356
HMKCZ06	H0392 H0488 H0675 L0040 L0439 L0764 L0771 L0773 L0774 L0775 S0360
HMEGG05	H0024 H0032 H0038 H0039 H0046 H0266 H0520 H0539 H0574 H0599 H0615
TIMEGGOS	H0622 H0632 H0633 L0021 L0163 L0439 L0483 L0598 L0608 L0646 L0654 L0655
	L0659 L0664 L0665 L0666 L0731 L0752 L0756 L0759 L0766 L0770 L0772 L0774
	L0775 L0776 L0783 L0809 S0003 S0007 S0038 S0116 S0142 S0194 S0214 S0356
	S0360 S0374 S0376 S0390 T0115
HCRNJ24	H0395 H0494 H0520 H0687 L0598 L0663 L0666 L0731 L0748 L0756 L0757 L0766
	L0770 L0775 L0792 L0794 L0803 S0356 S0380
HWLQA40	H0672 H0684 H0685 L0021 L0438 L0439 L0444 L0637 L0651 L0659 L0666 L0768
	L0769 L0771 L0772 L0774 L0775 S0360 S0376
HCQCC37	H0586 H0591 H0596 L0758 L0766 S0250 S0280 S0330 T0023
HACBB04	H0135 H0179 H0521 H0553 H0555 H0644 L0751 L0753 S0280
TIACDDV+	110155 110177 110521 110555 110555 110577 120755 20200

HWABJ67	H0063 H0479 H0581 L0521 L0592 L0659 L0665 L0666 L0759 L0776 S0360
HKLSD32	H0235 L0751 L0794
H2CAA34	L0773 L0779 T0103
H2CBN05	H0013 H0423 H0457 H0521 H0529 H0561 H0581 H0622 H0648 H0675 H0685
	H0702 L0105 L0366 L0655 L0658 L0659 L0665 L0667 L0748 L0758 L0762 L0766
	L0777 L0779 L0790 L0794 L0805 S0002 S0214 S0474 T0110
HCROZ66	S0356
HDPBY50	H0013 H0085 H0119 H0178 H0179 H0266 H0402 H0413 H0521 H0522 H0590
	H0593 H0596 H0622 H0650 H0656 H0657 H0672 L0021 L0143 L0364 L0382 L0483
	L0521 L0523 L0593 L0595 L0596 L0646 L0655 L0662 L0663 L0666 L0731 L0740
ŀ	L0744 L0745 L0747 L0750 L0752 L0754 L0755 L0761 L0764 L0766 L0775 L0789
	L0800 L0803 L0805 L0806 L0809 S0002 S0052 S0142 S0222 S0330 S0344 S0356
	S0358 S0360 S0362 S0374 S0378 S0380 S0472
HHPGT16	H0051 H0144 H0341 H0542 H0543 H0600 H0632 H0659 L0021 L0022 L0438 L0483
	L0663 L0666 L0731 L0744 L0755 L0758 L0759 L0766 L0774 L0777 L0783 S0038
HWAAD1	S0222 S0422 H0581 L0055 L0598 L0752 L0755 L0766 L0770 L0772 L0779 S0360
5 HWAADI	H0381 L0033 L0398 L0732 L0733 L0700 L0770 L0772 L0779 30300
HSPBY20	H0013 H0100 H0478 H0506 H0556 H0575 H0581 H0596 H0613 H0615 H0623
	H0647 L0361 L0438 L0471 L0483 L0485 L0591 L0599 L0608 L0740 L0747 L0758
	L0759 L0763 L0777 S0222 S3014 S6028
HCRQN90	H0081 S0356
HWBDR92	H0134 H0264 H0375 H0423 H0444 H0539 H0553 H0555 H0580 H0583 H0616
	H0667 L0369 L0662 L0738 L0754
HEMGF10	H0050 L0439 L0748 L0758 S0046
HWLXE78	H0392 L0372 L0541 L0604 L0659 L0731 L0775 L0777 L0794 L0803 S0360 T0040
***********	T0109
HCYBF60	L0362 L0748 T0114
HWLVY67	S0114 S0360
H2LAJ32	H0046 H0457 L0520 T0115 H0031 H0050 H0263 H0294 H0428 H0445 H0486 H0521 H0522 H0598 H0617
HDPQV66	H0031 H0030 H0203 H0294 H0428 H0443 H0480 H0321 H0322 H0398 H0017 H0661 H0664 H0667 H0688 L0439 L0471 L0483 L0518 L0602 L0608 L0642 L0659
	L0664 L0665 L0731 L0742 L0747 L0752 L0754 L0756 L0758 L0761 L0764 L0766
	L0770 L0771 L0776 L0779 L0800 L0805 L0809 S0003 S0010 S0049 S0134 S0330
	S0356
HBXFF23	H0393 H0438 H0661 L0740 S0376
HDPBG07	H0013 H0015 H0031 H0036 H0039 H0050 H0090 H0100 H0170 H0255 H0265
	H0271 H0305 H0306 H0352 H0402 H0423 H0424 H0435 H0445 H0486 H0494
	H0521 H0542 H0543 H0545 H0549 H0556 H0581 H0615 H0622 H0628 H0632
1	H0644 H0645 H0659 H0663 H0682 L0002 L0366 L0373 L0375 L0382 L0438 L0439
	L0483 L0517 L0527 L0532 L0564 L0588 L0592 L0599 L0601 L0638 L0663 L0731
	L0740 L0741 L0742 L0743 L0744 L0747 L0748 L0750 L0751 L0754 L0755 L0756
<i>.</i>	L0757 L0759 L0761 L0763 L0764 L0766 L0768 L0769 L0770 L0775 L0776 L0777
	L0783 L0794 L0800 L0805 L0806 L0809 S0002 S0007 S0029 S0036 S0049 S0126 S0134 S0140 S0192 S0346 S0354 S0360 S0376 S0420 T0010 T0042
HTDAI12	H0477
HWLGV14	L0748 S0354
HCYBM15	S0002 T0114
HTEPE28	H0038 H0369 H0616 L0593 L0595
HCQBI31	H0263 H0327 L0438 L0663 L0666 L0747 L0759 L0763 L0794 S0378
HCYBN59	H0051 T0114
HCRMK04	H0352 L0662 L0752 L0759 L0794 S0356
HE9RM22	H0009 H0100 H0144 H0333 L0352 L0438 L0512 L0651 L0748 L0749 L0759 L0776
	L0803 S0356 S0360
HJAAL27	H0591 L0749 L0758 L0767 L0775 L0794 S0358 S0376 S6028 T0041

HMUBP81	H0046 H0263 H0341 H0412 H0529 H0648 H0650 H0667 L0021 L0603 L0662 L0665			
	L0740 L0748 L0755 L0759 L0764 L0766 L0767 L0769 L0774 L0775 L0776 L0777			
	L0779 S0330 S0358 S0360 S0376 T0042			
НАРОТ58	H0575 L0731			
HCFLR18	H0423 H0485 L0651 S0150 S0356			
HDPAA38	H0521			
HWABG3	H0090 H0486 H0549 H0555 H0575 H0581 H0672 H0689 L0438 L0599 L0731 L0747			
2	L0748 L0755 L0758 L0768 L0770 L0775 L0779 L0780 L0788 L0794 S0002 S0142			
	S0144 S0344 S0356 S0360 S0376 S0462 S6014 T0041			
HKABL05	H0004 H0013 H0014 H0024 H0031 H0036 H0038 H0039 H0040 H0090 H0144			
	H0156 H0169 H0170 H0171 H0188 H0222 H0251 H0263 H0265 H0333 H0369			
	H0370 H0373 H0393 H0423 H0428 H0435 H0445 H0458 H0486 H0488 H0494			
	H0497 H0510 H0518 H0519 H0520 H0521 H0529 H0539 H0542 H0543 H0544			
	H0545 H0547 H0551 H0553 H0555 H0560 H0561 H0575 H0580 H0581 H0583			
	H0587 H0591 H0596 H0615 H0616 H0618 H0623 H0624 H0626 H0634 H0643			
	H0644 H0653 H0656 H0657 H0659 H0664 H0665 H0666 H0670 H0689 H0690			
	L0021 L0361 L0362 L0363 L0372 L0375 L0394 L0438 L0439 L0471 L0520 L0522			
	L0527 L0558 L0583 L0586 L0588 L0589 L0590 L0592 L0593 L0595 L0596 L0598			
	L0599 L0600 L0602 L0603 L0605 L0606 L0608 L0628 L0646 L0649 L0651 L0653			
	L0654 L0655 L0656 L0657 L0659 L0662 L0663 L0664 L0665 L0666 L0667 L0717			
	L0731 L0740 L0745 L0747 L0748 L0749 L0750 L0752 L0753 L0754 L0755 L0756			
	L0757 L0758 L0759 L0761 L0762 L0763 L0764 L0766 L0768 L0770 L0771 L0772			
	L0775 L0776 L0777 L0779 L0783 L0785 L0789 L0794 L0800 L0803 L0806 L0809			
	S0003 S0011 S0026 S0040 S0045 S0046 S0126 S0152 S0194 S0206 S0212 S0222			
	S0276 S0328 S0330 S0344 S0356 S0358 S0360 S0374 S0376 S0380 S0390 S0418			
TTE COMO	S0420 S0428 S0430 S0444 T0006 T0039 T0040 T0041 T0042 T0069			
HE8OT93	H0013 H0519 L0596 S0028 S0132 T0109			
HRODG74	H0156 H0455 H0598 H0619 H0659 L0021 L0040 L0439 L0638 L0750 L0754 L0756			
IID A DIZZZ	L0766 L0774 S0002 S0358 S0360 S0374 S0380			
HDABK73	H0263 H0497 L0601 H0009 H0136 H0435 H0494 H0587 H0598 H0634 L0759 S0132 S0242 S0332 S0448			
HOGDS65	T0110			
HHEGC16	H0542 H0543 H0551 H0575 L0731 L0740 L0745 L0748 L0766 L0770 S0126 S0360			
I HEGC10	S0420			
HETBI79	H0031 H0046 H0574 L0645 S0358 T0110			
HOENX50	H0052 H0097 H0144 H0170 H0171 H0263 H0318 H0341 H0373 H0412 H0413			
HOENASO	H0435 H0583 H0599 H0624 H0651 H0662 H0663 L0055 L0351 L0438 L0439 L0521			
	L0545 L0589 L0592 L0629 L0666 L0731 L0740 L0750 L0752 L0756 L0759 L0766			
	L0770 L0777 L0779 S0003 S0036 S0045 S0126 S0242 S6028			
HCEOW20	H0052 L0366 L0747 L0768 L0769 L0777 S0222 S6028			
НСЕРН79	H0052 H0619 L0748 L0750 L0756			
HKAEB15	H0052 H0135 H0159 H0265 H0266 H0494 H0521 H0529 H0539 H0542 H0553			
	H0560 H0575 H0599 H0677 L0456 L0596 L0601 L0764 S0027 S0040 S0045 S0192			
	S0356 S0380			
HCRMV17	H0069 L0761 S0356			
H2LBA37	H0013 H0624 S0356 T0115			
HWLIP86	H0014 H0052 H0596 L0764 L0772 L0773 L0775 S0356 S0358			
HOEMQ68	H0013 H0030 H0056 H0268 H0341 H0553 H0556 H0580 H0619 H0623 H0644			
I IIO E III QUU	L0748 S0011 S0028 S0126 S0150			
HUVHP60	H0039 H0040 H0050 H0216 H0264 H0328 H0445 H0506 H0510 H0519 H0542			
	H0547 H0551 H0560 H0594 H0623 H0625 H0638 H0657 L0439 L0598 L0648 L0662			
ļ	L0665 L0747 L0758 L0759 L0766 L0768 L0770 L0774 L0777 L0794 L0803 S0002			
	S0003 S0007 S0360 S0380 T0110			
HUFCI29	H0014 H0039 H0040 H0085 H0421 H0487 H0506 H0622 H0647 L0022 L0053 L0140			
]	L0194 L0364 L0372 L0373 L0483 L0596 L0599 L0627 L0640 L0659 L0662 L0667			
1	L0731 L0764 L0765 L0771 L0773 L0789 L0800 L0805 L0806 L0809 S0116 S0146			

	S0328 S0330 S0348 S0356 S0358 S0360 S0374 S0376 S0406 T0008 T0023
HAUAF56	H0294 H0341 S0010 S0031 S0053 T0060
HFCAI79	H0009 H0037 H0046 H0074 H0090 H0136 H0179 H0252 H0263 H0264 H0375
In Cattry	H0380 H0428 H0436 H0488 H0596 H0615 H0627 H0677 L0021 L0367 L0646 L0662
l	L0666 L0740 L0742 L0744 L0747 L0748 L0749 L0752 L0756 L0758 L0761 L0766
	L0777 L0779 L0803 L0806 S0116 S0122 S0280 S0356 S0358 S0360
HWHPZ02	H0012 H0046 H0100 H0266 H0352 H0421 H0428 H0478 H0484 H0485 H0494
	H0520 H0543 H0547 H0561 H0565 H0581 H0587 H0594 H0616 H0647 H0668
	H0670 H0689 L0105 L0372 L0471 L0475 L0637 L0665 L0698 L0731 L0744 L0747
	L0758 L0759 L0766 L0770 L0774 L0775 L0777 S0003 S0007 S0046 S0150 S0212
	S0280 S0354 S0356 S0376 S0422 S0424 S0460 T0041
HLTAZ90	H0090 L0780
HHFUM32	H0026 H0196 H0242 H0411 H0421 H0559 H0624 H0650 L0747 S0142
HHFAB62	H0019 L0439
HCEIC29	H0052 H0263 H0328 H0329 H0436 H0486 H0542 H0543 L0439 L0483 L0594 L0599
HEIGENICC	L0646 L0666 L0731 L0766 L0774 L0777 L0779 L0794 L0800 L0803 S0452 T0060
HFKFN66	H0012 L0748 H0023 H0032 H0039 H0040 H0046 H0056 H0083 H0132 H0144 H0207 H0214
HWMFQ1	H0216 H0520 H0521 H0553 H0555 H0590 H0591 H0619 L0005 L0143 L0595 L0608
"	L0740 L0742 L0745 L0750 S0027 S0142 S0150 S0152 S0196 S0212 S0328 S0354
	S0358
HCRBB01	H0012 H0031 H0036 H0038 H0039 H0040 H0046 H0050 H0059 H0063 H0081
	H0123 H0131 H0135 H0144 H0156 H0163 H0169 H0263 H0266 H0272 H0284
	H0286 H0309 H0318 H0327 H0341 H0345 H0370 H0375 H0380 H0391 H0411
	H0412 H0413 H0433 H0437 H0485 H0486 H0494 H0497 H0518 H0519 H0521
	H0539 H0545 H0546 H0547 H0553 H0581 H0583 H0586 H0587 H0593 H0594
	H0596 H0598 H0599 H0600 H0615 H0616 H0617 H0619 H0620 H0622 H0623
	H0628 H0638 H0644 H0645 H0646 H0647 H0648 H0652 H0653 H0658 H0659
	H0660 H0661 H0662 H0663 H0667 H0670 H0672 H0682 H0684 L0055 L0362 L0364 L0373 L0382 L0383 L0389 L0471 L0517 L0518 L0519 L0520 L0527 L0529 L0530
	L0542 L0595 L0596 L0598 L0602 L0631 L0646 L0648 L0651 L0657 L0659 L0664
	L0665 L0666 L0731 L0747 L0748 L0750 L0752 L0757 L0758 L0759 L0763 L0765
	L0768 L0769 L0771 L0774 L0775 L0776 L0777 L0783 L0806 S0002 S0003 S0010
	S0028 S0036 S0040 S0044 S0045 S0046 S0049 S0051 S0053 S0132 S0142 S0144
	S0148 S0150 S0192 S0194 S0208 S0214 S0278 S0306 S0320 S0322 S0328 S0330
1	S0334 S0344 S0354 S0356 S0358 S0360 S0370 S0372 S0374 S0376 S0378 S0382
	S0390 S0418 S0420 S0422 S0426 S0440 S0450 S0460 S0468 S6028 T0023 T0048
	T0067 T0115
HCROO59	H0036 L0644 L0758 S0114 S0356 T0006
HCRON34	S0356
HLQER45	H0014 H0015 H0037 H0040 H0204 H0316 H0431 H0506 H0509 H0517 H0574 H0596 H0597 H0598 H0622 H0632 H0642 H0675 L0363 L0372 L0382 L0581 L0596
	L0646 L0659 L0749 L0764 L0768 L0771 L0794 L0806 L0809 S0328 S0354 S0356
	S0358 S0360 S0374 S0376 S0434 S0456 S0472
HWLMK6	L0748 L0754 L0755 S0148 S0280 S0376 T0068 T0110
5	1 10/10 10/3/120/33 50110 50200 503/0 10000 10/10
HNTBD04	H0009 H0014 H0050 H0063 H0208 H0225 H0352 H0413 H0445 H0459 H0494
	H0497 H0519 H0520 H0521 H0591 H0632 H0646 H0656 H0657 H0663 H0667
	L0364 L0483 L0581 L0663 L0744 L0748 L0752 L0761 L0766 L0770 L0777 L0779
	L0800 S0002 S0010 S0026 S0028 S0210 S0212 S0346 S0356 S0358 S0376 S0420
	T0002
HSUSF13	H0046 H0144 H0163 H0216 H0251 H0316 H0412 H0428 H0435 H0486 H0494
	H0519 H0520 H0529 H0539 H0561 H0587 H0624 H0628 H0634 H0660 H0672
	L0731 L0740 L0745 L0747 L0750 L0752 L0754 L0757 L0758 S0028 S0045 S0049
IIIIVEC16	S0126 S0192 S0206 S0212 S0242 S0360 S0436
HUVFS16	H0046 H0343 H0521 H0560 H0623 L0532 L0777 S0028 S0046 S0356 S0386

HCQBD51	H0263 H0648 H0658 L0362 L0527 L0663 L0664 L0740 L0758 L0766 L0774 L0804
HCRNO79	H0090 S0356
HCRND67	S0356
HAVTF85	S0328 S0414
HWLVS52	S0360
HCRNJ46	H0046 S0356
HCRPE30	S0356
HKGAW0	L0375
2	
HWLOW7	H0316 H0351 H0423 H0435 H0441 H0581 H0616 H0622 H0658 L0021 L0438 L0439
2	L0521 L0591 L0650 L0657 L0662 L0663 L0666 L0744 L0745 L0747 L0748 L0749
	L0754 L0755 L0759 L0764 L0768 L0769 L0775 L0777 L0779 L0788 L0794 L0804
	S0026 S0356 S0376
HUSGT72	H0309 H0412 H0422 L0745 S0360
HPWBM9	H0156 H0615 H0658 L0526 L0662 L0663 L0758 L0759 L0761 L0763 L0766 L0768
1	L0769 L0770 L0809 S0007 S0044 S0360
HWLRC68	S0132 S0360
HKAOG63	H0032 H0040 H0046 H0071 H0081 H0085 H0134 H0144 H0150 H0169 H0265
	H0284 H0428 H0445 H0494 H0506 H0521 H0556 H0574 H0599 H0624 H0659
	L0439 L0471 L0518 L0588 L0645 L0662 L0663 L0666 L0731 L0740 L0743 L0744
	L0747 L0748 L0749 L0750 L0752 L0754 L0755 L0758 L0759 L0763 L0764 L0766
	L0767 L0768 L0769 L0770 L0771 L0774 L0775 L0776 L0777 L0780 L0804 L0805
	S0027 S0028 S0036 S0049 S0146 S0152 S0212 S0214 S0278 S0358 S0360 S0374
	S0376 S0420 S6028 T0042 T0048 T0049
H2LAC34	H0123 H0419 H0547 H0553 L0055 L0387 L0438 L0439 L0649 L0657 L0731 L0748
	L0751 L0752 L0757 L0759 L0766 L0768 L0769 L0773 L0774 L0775 L0776 L0779 L0789 L0806 S0046 S0360 S0424 T0010 T0041 T0115
THUED A 07	
HHFBA07	H0050 S0356
HWHGC9	H0586 L0744
HMCDH54	H0007 H0052 H0255 H0580 H0616 H0687 L0646 L0752 L0758 S0002 S0037 S0142
HEGAM94	H0024 H0038 H0046 H0050 H0052 H0090 H0101 H0263 H0370 H0445 H0455
IIEGAW54	H0550 H0555 H0556 H0633 H0641 H0667 H0672 L0055 L0352 L0438 L0439 L0517
	L0520 L0630 L0659 L0662 L0666 L0740 L0743 L0747 L0748 L0749 L0750 L0751
	L0752 L0754 L0758 L0759 L0763 L0766 L0775 L0779 S0003 S0052 S0116 S0126
	S0192 S0196 S0250 S0276 S0356 S0360 S0376
HARAG42	H0231 H0263 H0597 L0764 S0358 S0374 S0376 T0082
HCQDL20	H0013 H0036 H0090 H0144 H0393 H0587 H0590 H0596 H0622 H0632 L0662
	L0748 S0356
HLQGF34	H0632 S0356
HMIBE59	L0754 S0044 S0126 S6028
H6EDF71	H0083 H0090 H0222 H0265 H0370 H0423 H0445 H0486 H0506 H0519 H0529
	H0543 H0553 H0559 H0581 H0624 H0638 H0648 H0658 H0667 H0672 H0674
	H0685 L0366 L0475 L0521 L0599 L0646 L0651 L0663 L0665 L0666 L0747 L0750
	L0755 L0766 L0775 L0777 L0783 L0788 S0002 S0003 S0044 S0152 S0276 S0354
	S0360 S0374 S0376
HHFMH12	H0008 H0014 H0023 H0040 H0048 H0050 H0052 H0083 H0100 H0194 H0231
	H0263 H0266 H0272 H0295 H0321 H0370 H0393 H0412 H0413 H0433 H0435
	H0494 H0519 H0520 H0543 H0556 H0619 H0622 H0623 H0635 H0665 H0691
	L0157 L0366 L0483 L0601 L0655 L0659 L0662 L0663 L0717 L0731 L0742 L0748
	L0749 L0753 L0756 L0758 L0759 L0769 L0770 L0777 L0803 L0809 S0027 S0036
	S0038 S0044 S0046 S0114 S0132 S0134 S0144 S0194 S0210 S0222 S0242 S0250
	S0306 S0356 S0360 S0380 S3014 T0010 T0040 T0041 T0048
HCFMY07	H0040 H0046 H0170 H0271 H0422 H0423 H0521 H0543 H0555 H0572 H0583
	H0650 L0371 L0384 L0462 L0471 L0475 L0483 L0592 L0595 L0598 L0599 L0603

	L0666 L0740 L0748 L0758 L0759 L0764 L0777 L0779 L0794 S0003 S0126 S0222
HOVDDAG	S0242 S0250 S0360 S0364 S0418
HSYBP46	H0008 H0013 H0014 H0019 H0024 H0031 H0038 H0039 H0050 H0052 H0057
	H0090 H0123 H0144 H0170 H0171 H0197 H0244 H0251 H0266 H0268 H0328
1	H0329 H0370 H0393 H0412 H0427 H0485 H0486 H0519 H0520 H0528 H0544
	H0551 H0574 H0586 H0591 H0593 H0595 H0598 H0619 H0622 H0624 H0631
	H0644 H0646 H0653 H0659 H0665 H0668 L0021 L0373 L0375 L0389 L0471 L0564
ŀ	L0565 L0590 L0591 L0598 L0603 L0606 L0646 L0659 L0662 L0665 L0666 L0731
	L0738 L0740 L0743 L0744 L0747 L0748 L0749 L0750 L0752 L0754 L0757 L0758
	L0759 L0763 L0771 L0777 L0779 L0784 L0794 L0806 S0001 S0003 S0011 S0013
	S0022 S0026 S0028 S0032 S0040 S0046 S0126 S0192 S0194 S0206 S0210 S0212
	S0214 S0250 S0342 S0356 S0360 S0374 S0376 S0380 S0384 S0418 S0420 S3014
	T0039 T0049 T0114
HE8DZ94	H0013 H0031 H0038 H0130 H0251 H0263 H0266 H0268 H0428 H0435 H0445
	H0486 H0509 H0522 H0535 H0539 H0543 H0575 H0586 H0598 H0616 H0644
	H0657 H0662 H0666 H0670 H0672 H0674 L0021 L0371 L0372 L0523 L0526 L0542
	L0596 L0603 L0637 L0662 L0663 L0740 L0745 L0746 L0748 L0750 L0755 L0758
	L0759 L0764 L0766 L0768 L0770 L0774 L0775 L0779 L0803 L0809 S0002 S0003
VID OD DOG	S0010 S0044 S0046 S0126 S0214 S0222 S0356 S0358 S0360 S0380 T0049
HRGDE77	H0014 H0040 H0052 H0102 H0123 H0134 H0144 H0242 H0399 H0457 H0483
	H0622 H0624 H0689 L0605 L0748 L0751 S0002 S0010 S0045 S0144 S0218 S0222
VI	S0278 S0344 S0356 S0360
HMUBQ39	H0036 H0038 H0052 H0083 H0123 H0131 H0156 H0272 H0309 H0318 H0369
	H0384 H0393 H0413 H0422 H0478 H0494 H0506 H0509 H0520 H0522 H0529
	H0538 H0551 H0555 H0587 H0624 H0634 H0638 H0644 H0657 H0663 H0670
	H0672 L0055 L0352 L0362 L0369 L0383 L0435 L0438 L0439 L0471 L0540 L0590
	L0591 L0594 L0595 L0596 L0605 L0608 L0646 L0647 L0652 L0659 L0663 L0665
	L0740 L0747 L0749 L0754 L0758 L0759 L0761 L0764 L0766 L0770 L0777 L0794
	L0803 L0804 L0806 S0002 S0026 S0028 S0037 S0150 S0192 S0194 S0206 S0342 S0354 S0356 S0358 S0360 S0420 S0422 S6026 T0039 T0067 T0071
HCEYN60	H0030 H0038 H0052 H0087 H0097 H0116 H0153 H0253 H0261 H0265 H0295
I ICE I NOU	H0309 H0327 H0333 H0422 H0423 H0483 H0486 H0506 H0519 H0521 H0545
ļ	H0546 H0593 H0616 H0618 H0620 H0647 H0651 L0055 L0369 L0375 L0383 L0384
	L0394 L0439 L0503 L0509 L0510 L0511 L0581 L0588 L0592 L0593 L0596 L0600
	L0634 L0655 L0666 L0731 L0742 L0743 L0744 L0748 L0749 L0751 L0754 L0757
	L0758 L0764 L0766 L0768 L0769 L0770 L0772 L0776 L0785 L0809 S0132 S0142
	S0144 S0278 S0356 S0360 S0376 S0418 S0420 T0002 T0049 T0082
HTWEA61	H0436
HLSAA96	H0040 H0046 H0051 H0052 H0123 H0144 H0156 H0178 H0204 H0239 H0253
IILSAA70	H0265 H0423 H0488 H0539 H0540 H0549 H0556 H0596 H0617 H0635 H0658
ļ.	L0369 L0375 L0439 L0530 L0648 L0659 L0663 L0665 L0747 L0748 L0750 L0758
	L0759 L0766 L0770 L0777 L0779 L0803 S0002 S0028 S0142 S0410
HTEEZ62	H0031 H0038 H0039 H0046 H0050 H0144 H0271 H0318 H0329 H0331 H0412
11122202	H0423 H0428 H0431 H0435 H0436 H0441 H0445 H0478 H0486 H0488 H0494
	H0497 H0506 H0519 H0520 H0521 H0543 H0565 H0572 H0581 H0586 H0615
ì	H0616 H0624 H0627 H0631 H0648 H0650 H0657 H0660 H0670 H0672 H0682
	H0687 L0364 L0372 L0438 L0471 L0485 L0521 L0523 L0527 L0591 L0592 L0599
	L0655 L0659 L0662 L0663 L0664 L0665 L0666 L0740 L0747 L0748 L0749 L0752
	L0755 L0756 L0758 L0763 L0764 L0770 L0772 L0774 L0776 L0777 L0787 L0794
	L0800 L0803 L0804 L0805 L0807 L0809 S0003 S0007 S0027 S0036 S0044 S0045
	S0046 S0053 S0114 S0116 S0126 S0144 S0192 S0278 S0280 S0328 S0330 S0342
1	S0354 S0356 S0358 S0360 S0374 S0378 S0422 S0424 S0428 S0436 S0444 S0446
<b>\</b>	S0462 S0474 T0041 T0109 T0110
HOAAH52	H0031 H0032 H0038 H0046 H0052 H0056 H0057 H0059 H0090 H0105 H0135
	H0144 H0169 H0252 H0316 H0318 H0328 H0331 H0369 H0381 H0392 H0412
	H0423 H0431 H0435 H0457 H0486 H0506 H0521 H0522 H0543 H0545 H0546
·	

Г	H0553 H0555 H0556 H0560 H0561 H0563 H0569 H0574 H0581 H0586 H0591
	H0553 H0553 H0556 H0560 H0561 H0563 H0569 H0574 H0581 H0586 H0591
1	H0664 H0672 L0352 L0369 L0378 L0438 L0439 L0518 L0521 L0565 L0581 L0595
1	L0596 L0598 L0605 L0646 L0656 L0659 L0662 L0663 L0664 L0666 L0717 L0731
1	L0740 L0742 L0747 L0748 L0749 L0750 L0752 L0754 L0755 L0757 L0758 L0762
	L0764 L0770 L0771 L0773 L0774 L0775 L0776 L0777 L0794 L0796 L0803 L0806
	S0002 S0007 S0010 S0026 S0027 S0044 S0112 S0114 S0146 S0212 S0222 S0242
	S0250 S0314 S0364 S0380 S0394 S0424 S0444 S3014
HFKMJ24	H0046 H0271 H0542 H0543 H0620 L0738 S0360
HEOQC11	H0024 H0030 H0032 H0038 H0040 H0051 H0087 H0100 H0123 H0135 H0165
	H0181 H0200 H0264 H0266 H0271 H0290 H0294 H0295 H0316 H0318 H0369
	H0370 H0422 H0424 H0427 H0428 H0457 H0510 H0539 H0543 H0556 H0581
1	H0597 H0616 H0618 H0619 H0634 H0644 H0657 H0667 H0684 L0372 L0528 L0587
1	L0593 L0596 L0599 L0617 L0627 L0646 L0658 L0659 L0662 L0665 L0731 L0748
	L0749 L0751 L0752 L0756 L0759 L0764 L0766 L0768 L0770 L0775 L0783 L0809
	S0027 S0037 S0126 S0134 S0152 S0192 S0354 S0360 S0376 S0418 T0039
HCEDM42	H0009 H0040 H0052 H0056 H0083 H0087 H0169 H0264 H0265 H0333 H0413
	Н0484 Н0486 Н0497 Н0543 Н0556 Н0581 Н0592 Н0597 Н0598 Н0617 Н0620
	H0646 H0656 H0657 L0439 L0588 L0601 L0655 L0663 L0731 L0740 L0747 L0748
	L0763 L0764 L0766 L0769 L0774 L0779 L0780 S0046 S0126 S0142 S0196 S0242
	S0278 S0356 S0456 T0049 T0110
HWMBU8	H0030 S0376
9	
HMKAN7	H0032 H0156 H0251 H0375 H0392 H0436 H0547 H0561 H0600 H0616 H0634
1	H0638 H0657 H0659 L0388 L0438 L0518 L0581 L0588 L0605 L0649 L0655 L0659
	L0661 L0662 L0663 L0665 L0666 L0717 L0731 L0748 L0754 L0758 L0759 L0766
	L0768 L0773 L0775 L0779 L0794 L0803 L0806 S0003 S0026 S0037 S0053 S0356
	S0360 S0422
HSIFV30	H0013 H0030 H0032 H0036 H0050 H0056 H0083 H0085 H0124 H0130 H0170
	H0171 H0213 H0222 H0231 H0232 H0247 H0253 H0263 H0265 H0266 H0318
	H0373 H0403 H0412 H0423 H0424 H0506 H0521 H0542 H0551 H0556 H0575
	H0587 H0590 H0592 H0597 H0599 H0634 H0646 H0660 L0040 L0438 L0439 L0564
	L0592 L0596 L0627 L0655 L0659 L0665 L0717 L0731 L0740 L0748 L0750 L0751
	L0758 L0759 L0764 L0765 L0766 L0767 L0768 L0770 L0771 L0774 L0777 L0794
	L0804 L0805 S0028 S0344 S0346 S0354 S0356 S0358 S0364 S0374 S0376 S0386
	T0006 T0049
HNTSY52	H0023 H0032 H0038 H0039 H0042 H0051 H0083 H0090 H0171 H0196 H0251
	H0266 H0329 H0373 H0412 H0423 H0477 H0478 H0509 H0521 H0543 H0547
	H0553 H0575 H0580 H0581 H0591 H0595 H0644 H0659 H0667 L0471 L0485 L0512
	L0588 L0608 L0637 L0646 L0647 L0650 L0653 L0659 L0661 L0662 L0663 L0666
	L0731 L0740 L0745 L0750 L0754 L0755 L0756 L0757 L0759 L0761 L0763 L0765
	L0766 L0770 L0771 L0803 L0806 S0003 S0026 S0028 S0048 S0112 S0126 S0134
	S0152 S0192 S0242 S0250 S0260 S0282 S0328 S0354 S0356 S0358 S0360 S0374
	S0376 S0422 S0424 T0041 T0067 T0115
HWLKD85	H0009 H0013 H0014 H0030 H0038 H0039 H0040 H0046 H0052 H0090 H0150
	H0171 H0251 H0252 H0264 H0265 H0266 H0295 H0318 H0355 H0375 H0393
	H0422 H0438 H0458 H0478 H0484 H0494 H0521 H0529 H0543 H0544 H0547
	H0551 H0560 H0575 H0581 H0615 H0625 H0635 H0647 H0653 H0657 H0661
	H0673 H0686 H0693 L0021 L0163 L0364 L0366 L0382 L0500 L0521 L0526 L0564
	L0599 L0601 L0608 L0637 L0638 L0649 L0659 L0662 L0665 L0666 L0745 L0747
	L0749 L0750 L0752 L0754 L0755 L0759 L0764 L0766 L0767 L0768 L0769 L0772
	L0773 L0775 L0777 L0779 L0780 L0794 L0796 L0809 S0031 S0040 S0114 S0126
	1 S0192 S0210 S0242 S0328 S0330 S0356 S0358 S0360 S0376 S0380 S0386 S0418
	S0192 S0210 S0242 S0328 S0330 S0356 S0358 S0360 S0376 S0380 S0386 S0418
HWI EI 48	S3012
HWLEL48 HADME31	

HFVJL45	H0331 H0393 H0510 L0748 S0440
HLJEA63	H0375 S0354
HAIBW90 H2CBE03	H0024 H0032 H0038 H0039 H0040 H0063 H0100 H0130 H0169 H0170 H0264 H0328 H0331 H0333 H0343 H0352 H0369 H0412 H0413 H0428 H0436 H0484 H0486 H0494 H0506 H0520 H0544 H0545 H0553 H0574 H0575 H0581 H0587 H0591 H0594 H0595 H0616 H0619 H0622 H0624 H0628 H0648 H0657 H0671 H0684 L0005 L0065 L0362 L0369 L0375 L0383 L0438 L0485 L0515 L0521 L0598 L0600 L0608 L0623 L0637 L0638 L0647 L0656 L0659 L0662 L0663 L0664 L0665 L0666 L0698 L0731 L0743 L0745 L0749 L0750 L0751 L0754 L0755 L0756 L0757 L0758 L0763 L0766 L0768 L0769 L0770 L0774 L0776 L0777 L0779 L0796 L0803 L0805 L0806 S0002 S0003 S0007 S0010 S0027 S0028 S0051 S0126 S0132 S0134 S0194 S0206 S0212 S0214 S0222 S0260 S0280 S0328 S0350 S0358 S0360 S0414 S0422 S0426 S3014 S6026 T0006 T0039 T0048 T0049 T0109
HSRBB92	H0013 H0024 H0040 H0059 H0124 H0144 H0208 H0334 H0341 H0370 H0421
nsidb <sub>2</sub>	H0438 H0520 H0521 H0543 H0547 H0551 H0592 H0596 H0597 H0599 H0616 H0617 H0650 H0653 H0658 H0670 H0694 L0163 L0362 L0438 L0439 L0455 L0456 L0592 L0599 L0608 L0649 L0659 L0662 L0731 L0740 L0747 L0748 L0749 L0751 L0754 L0758 L0759 L0761 L0763 L0764 L0766 L0769 L0770 L0771 L0772 L0774 L0776 L0779 L0780 L0785 L0789 L0806 S0011 S0132 S0144 S0328 S0360 S0366 S0378 S0386 T0042 T0110
HSYEA10	H0039 H0040 H0056 H0069 H0179 H0200 H0250 H0268 H0269 H0372 H0412 H0413 H0433 H0437 H0486 H0551 H0593 H0617 H0622 H0623 H0628 H0647 L0021 L0376 L0483 L0565 L0588 L0602 L0646 L0647 L0648 L0653 L0659 L0662 L0757 L0759 L0764 L0794 L0806 S0036 S0045 S0046 S0049 S0132 S0196 S0242
	S0332 S0350 S0356 S0360 S0376 S0450 S0468 T0008 T0039 T0049
HNTAR08	H0012 H0014 H0030 H0037 H0040 H0123 H0124 H0132 H0144 H0176 H0179 H0187 H0194 H0250 H0252 H0253 H0271 H0290 H0295 H0309 H0370 H0402 H0411 H0412 H0416 H0427 H0519 H0539 H0543 H0545 H0546 H0549 H0553 H0586 H0592 H0618 H0619 H0620 H0622 H0675 H0684 H0685 H0687 L0021 L0142 L0163 L0364 L0376 L0383 L0438 L0439 L0471 L0591 L0593 L0595 L0603 L0608 L0617 L0640 L0644 L0646 L0655 L0659 L0662 L0663 L0665 L0666 L0717 L0731 L0740 L0741 L0748 L0750 L0751 L0754 L0756 L0766 L0767 L0768 L0769 L0772 L0774 L0776 L0777 L0794 L0803 L0809 S0002 S0038 S0052 S0053 S0132 S0210 S0216 S0250 S0306 S0330 S0364 S0366 S0426 S6028 T0115
HNGEF72	S0052
HDHEA53	H0013 H0570
HCHAC08	H0024 H0051 H0170 H0483 H0494 H0576 H0617 H0622 H0657 L0596 L0603 L0749 S0354 S0358 S0376 S3014
HTLEN01	H0009 H0024 H0069 H0087 H0123 H0253 H0254 H0264 H0351 H0370 H0428 H0484 H0488 H0494 H0520 H0521 H0538 H0543 H0593 H0618 H0684 H0687 L0439 L0534 L0539 L0562 L0741 L0758 L0766 S0344 S0356 S0360
HCROA43	\$0356
HBINP81	H0046 H0494 H0593 H0622 S0356
HDLAG89	H0012 H0024 H0031 H0038 H0171 H0252 H0411 H0431 H0457 H0484 H0543 H0581 H0583 H0616 H0622 H0633 L0366 L0471 L0592 L0662 L0740 L0748 L0759 L0766 L0774 S0036 S0146 S0192 S0210 S0330 S0360 S0376 T0006
HWLDZ74	H0036 H0046 H0136 H0144 L0626 L0754 S0027 S0028 S0045 S0312 S0374
HNTST71 HNDAD16	H0039 H0547 H0622 S0358 H0046 H0081 H0156 H0231 H0483 H0484 H0517 H0597 H0617 H0670 L0666 L0747 L0762 L0764 L0771 L0790 L0800 S0358 S0360 S0374 S0376 T0109
HSVCD79	H0309 L0754 L0758 S0011 S0192 S0356
HCQDE22	H0596 L0749
HWLVU33	L0758 S0360
HKGBP52	H0040 H0331 H0538 L0005 L0351 L0439 L0731 L0740 L0748 L0749 L0754 L0757 L0758 L0775 L0776 L0800 S0356 S0360 S0426

	70.000 X 0.000 X 0.000 X 0.000 X 0.000 X 0.000 X 0.0000 X 0.0000 X 0.0000			
HDPPB40	H0522 L0438 L0439 L0662 L0748 L0752 L0758 L0777 L0803			
HCRMJ47	H0086 H0352 H0544 H0545 H0553 S0356			
HCRNK75	H0046 H0051 H0059 H0071 H0355 H0428 H0438 H0479 H0510 H0583 H0615			
	H0644 H0660 H0670 H0672 L0142 L0366 L0604 L0751 L0763 L0775 L0777 S0026			
	S0028 S0031 S0192 S0282 S0356 S0424			
HOGDR01	H0264 H0435 H0672 L0659 L0662 L0666 L0779 L0803 S0260 S0328 S0376 S0472			
HDPBW68	H0040 H0046 H0255 H0264 H0423 H0497 H0518 H0521 H0556 H0580 H0586			
	H0618 H0620 H0641 H0656 H0682 L0590 L0657 L0659 L0748 L0749 L0751 L0758			
	L0761 L0764 L0768 L0770 L0776 L0777 L0785 L0794 L0805 S0212 S0380 S0460			
HNTRV11	H0004 H0014 H0030 H0031 H0032 H0039 H0124 H0144 H0171 H0268 H0341			
	H0375 H0411 H0412 H0416 H0427 H0428 H0445 H0497 H0506 H0539 H0544			
	H0547 H0551 H0553 H0575 H0581 H0586 H0592 H0593 H0596 H0600 H0616			
	H0618 H0622 H0623 H0633 H0643 H0644 H0646 H0647 H0648 H0659 H0662			
	H0670 H0687 L0438 L0439 L0512 L0591 L0595 L0596 L0599 L0600 L0606 L0646			
	L0655 L0657 L0659 L0662 L0663 L0664 L0666 L0731 L0740 L0744 L0748 L0749			
,	L0750 L0752 L0754 L0755 L0756 L0757 L0758 L0761 L0774 L0775 L0776 L0777			
	L0779 L0785 L0800 L0805 S0003 S0040 S0045 S0046 S0052 S0176 S0192 S0196			
	S0222 S0242 S0276 S0314 S0318 S0354 S0356 S0358 S0360 S0370 S0374 S0376			
	S0388 S0392 S0454 T0110			
HBSAK60	H0204 H0205 H0381 H0624 S0028 S0044 S0050 S0053 S0260 S0282 T0074			
HDPOD73	H0521 H0522 H0581 L0021 L0759 S0356 S0360			
HWLKM7	H0036 H0050 H0486 H0593 S0358			
7	110030 110030 110400 110393 50330			
HCRMU04	H0012 H0014 H0031 H0038 H0040 H0050 H0087 H0123 H0144 H0250 H0352			
HCKMU04	H0373 H0413 H0435 H0478 H0494 H0521 H0522 H0529 H0556 H0586 H0620			
	H0625 H0632 H0644 H0646 H0650 H0658 H0682 L0157 L0365 L0369 L0438 L0439			
	L0471 L0512 L0565 L0599 L0638 L0642 L0645 L0647 L0659 L0662 L0665 L0666			
	L0717 L0740 L0747 L0748 L0749 L0750 L0751 L0753 L0754 L0757 L0758 L0759			
	L0763 L0764 L0766 L0768 L0769 L0770 L0771 L0774 L0777 L0779 L0789 L0794			
	L0806 L0809 S0001 S0002 S0003 S0010 S0011 S0036 S0037 S0040 S0052 S0126			
	S0132 S0192 S0206 S0216 S0222 S0278 S0280 S0294 S0354 S0356 S0360 S0418			
******	S0420 S3014 T0114			
HSSGC06	H0002 H0012 H0022 H0024 H0038 H0046 H0050 H0068 H0097 H0134 H0135			
	H0163 H0265 H0286 H0341 H0416 H0427 H0457 H0521 H0529 H0547 H0551			
	H0553 H0575 H0581 H0587 H0591 H0592 H0596 H0616 H0619 H0641 H0658			
	H0687 H0696 L0375 L0591 L0596 L0599 L0601 L0611 L0659 L0664 L0665 L0731			
	L0748 L0749 L0750 L0756 L0757 L0758 L0764 L0766 L0769 L0770 L0772 L0775			
	L0780 L0786 L0805 L0809 S0132 S0134 S0150 S0152 S0182 S0218 S0278 S0332			
	S0358 T0042 T0114			
HWLMO7	H0445 S0376			
3 H6ED 104	H0257			
H6EBJ04	H0257			
HCNDF58	H0597 S0358 L0608 S0010 S0376			
HWMBS0				
8	H0031 H0052 H0436 H0519 H0619 H0672 L0438 L0439 L0456 L0740 L0742 L0743			
HCRQL67				
	L0747 L0748 L0749 L0750 L0754 L0755 L0759 L0768 L0769 L0770 L0779 L0794			
WOD) 10.55	S0001 S0007 S0354 S0356 S0358 S0408 T0010			
HCRMO57	H0038 H0039 H0042 H0059 H0063 H0165 H0292 H0328 H0370 H0431 H0435			
	H0484 H0494 H0506 H0555 H0593 H0615 H0616 H0647 H0660 H0672 H0673			
	H0689 L0517 L0518 L0596 L0600 L0601 L0666 L0758 L0789 L0809 S0015 S0044			
	S0354 S0358 S0360 S0374 S0376 S0432			
HMUBO53	H0013 H0014 H0036 H0040 H0050 H0083 H0135 H0144 H0171 H0231 H0251			
	H0305 H0309 H0318 H0341 H0376 H0402 H0412 H0413 H0422 H0423 H0428			
	H0436 H0444 H0445 H0486 H0510 H0512 H0519 H0520 H0521 H0529 H0543			
	H0551 H0555 H0560 H0574 H0576 H0581 H0583 H0617 H0624 H0648 H0653			
	110551 110561 110561 110561 110561 110017 110024 110046 110033			

	H0656 H0657 H0658 H0659 H0663 H0666 H0674 H0684 L0021 L0362 L0363 L0367
	L0369 L0372 L0373 L0375 L0389 L0518 L0520 L0521 L0526 L0529 L0553 L0588
	L0592 L0593 L0595 L0596 L0599 L0608 L0636 L0637 L0648 L0662 L0663 L0664
	L0665 L0666 L0667 L0731 L0747 L0758 L0764 L0766 L0768 L0774 L0776 L0777
	L0804 L0805 L0809 S0003 S0026 S0050 S0116 S0328 S0330 S0358 S0360 S0374
	S0380 S0424 S3014 S6024 T0067
HDPMJ48	H0013 H0144 H0169 H0171 H0186 H0196 H0318 H0341 H0392 H0411 H0412
	H0483 H0510 H0521 H0522 H0551 H0561 H0581 H0596 H0615 H0624 H0634
	H0647 H0670 L0646 L0657 L0666 L0731 L0747 L0766 L0769 L0772 S0046 S0356
	S0358 S0360 T0041
HCQCD92	H0004 H0014 H0030 H0031 H0032 H0039 H0124 H0144 H0171 H0268 H0341
	H0375 H0411 H0412 H0416 H0427 H0428 H0445 H0497 H0506 H0539 H0544
	H0547 H0551 H0553 H0556 H0575 H0581 H0586 H0592 H0593 H0596 H0600
	H0616 H0618 H0622 H0623 H0633 H0643 H0644 H0646 H0647 H0648 H0659
	H0662 H0670 H0687 L0438 L0439 L0512 L0591 L0595 L0596 L0599 L0600 L0606
	L0646 L0655 L0657 L0659 L0662 L0663 L0664 L0666 L0731 L0740 L0744 L0748
	L0749 L0750 L0752 L0754 L0755 L0756 L0757 L0758 L0761 L0774 L0775 L0776
	L0777 L0779 L0785 L0800 L0805 S0003 S0040 S0045 S0046 S0052 S0152 S0176
	S0192 S0196 S0222 S0242 S0276 S0314 S0318 S0354 S0356 S0358 S0360 S0370
	S0374 S0376 S0388 S0392 S0454 T0110
HCQDS58	H0263 H0586 H0596 H0657 L0665 L0756 L0777
HCROE42	H0596 L0805 S0350 S0356
HCRMW6	H0263 H0313 H0428 H0445 H0486 H0539 H0667 L0471 L0594 L0758 L0766 L0777
2	S0026 S0194 S0356
HCRNR03	L0438 L0439 L0752 L0766 S0356
HWMCK5	H0188 H0660 H0663 L0021 L0775 L0803 S0282 S0376 S0424
1	

Table 4

SEQ ID	Cytologic Band or	OMIM Reference(s):
NO: X	Chromosome:	
3	10q26.13-q26.3	263700
5	11p15.1	114130 168450 257200 276904 600509 600937 601387 602092
9	6q21-q22	120110 121014 156225 164200 601410 601666 601757 602772
17	11q22	105580 133780 602574
18	3p21.3	116806 120120 120436 138320 168468 182280 600163
26	Xp11.21	300047 301300 301830 305400 308300 309470 309500 309610 311050
35	18	
36	Xq13	300011 300032 301310 305100 305450 308380 309605 311800 311870 314580
40	22q11.23	123620 600850
42	11q22-q23	105580 107680 107720 133780 147791 159555 168000
		186740 186830 188025 203750 208900 261640 600048 601382 602574
46	17q12-q21	109270 113705 144200 148065 148066 148067 148069
		148080 154275 168610 171190 176705 180240 182138
		185800 200350 221820 232200 249000 252920 253250
		600119 600881 601363 601687 601844 601954
47	19q13.2	107741 113900 122720 126340 126391 160900 164731 173850 207750 248600 258501
64	7q22	126650 154276 173360 602136 602447
69	12p12.1	112410 150100 168470 190070 200990 602096
76	22q12.2	101000 123620 138981 188826 600850 601669
81	5q11.2-q13.1	126060 143200 181510 214300 253200 268800 600354 600887
85	14q24.3-q31	104311 109150 182600 245200 275200 601208
92	2p22	120435 182601 278300 601071 601771 602134
103	21q22.3	120220 120240 123580 151385 171860 190685 236100 236200 240300 267750 600065 601072 601145
123	6p21.3	106300 108800 120290 120810 120820 142857 142858
	•	150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261 601868 602280 602475
139	6q27	152200 167000 600320 600883 602544
150	1p35.1	118210 120550 120570 120575 121800 130500 133200 171760 185470 230350 255800 602771
151	10	
163	2p21	120435 126600 135300 136435 152790 157170 182601 601771
167	12q24.3	160781 181405
173	7q33	180105 222800
174	22q12.3	138981 188826 190040 600850 601669
177	22q13.33	
192	3q23	106165 110100 117700 150210 169600 180380 203500

		276902 601199 601682
194	3p24	227646 600163 601154
196	Xq25	300123 301201 301845 304340 307150 308240 310490
190	Aq23	312000 313850
197	18	012000 012000
207	16p12-p11.2	108730 147781 172471 182381 186580 266600 600760
_0,	107127111	600761 602066
210	1p34.1-p32	120260 120550 120570 120575 120950 120960 121800
		130500 133200 138140 168360 171760 176100 178300
		187040 230000 246450 255800 600101 600650 600722
212	9q33.1	268900
216	14q32.33	144120 147020 147110
217	22q12.2	101000 123620 138981 188826 600850 601669
221	5q13	126060 143200 181510 253200 268800 600354
227	20q13	600281
228	Xq21.3-q22	300088 300300 301201 301500 301835 303400 303630
		303631 304500 304700 305450 309300 309605 311850
		312080
235	16p12	108730 147781 172471 186580 266600 600760 600761
		602066
242	19p13.2	108725 120700 133171 143890 147670 151440 164953
		231670 600276 600957 601843
244	6p12	180297 230450 263200 601690
254	6p21.3,	106300 108800 120290 120810 120820 142857 142858
		150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
255	15q22.1	151670 191010 601780
256	12q24.1	124200 147440 160781 181405 261600 601406 601620
		601621
258	17q21	109270 113705 144200 148065 148066 148067 148069
		148080 154275 168610 171190 176705 185800 200350
		221820 232200 249000 252920 253250 600119 601363
		601844
259	19q13.2	107741 113900 122720 126340 126391 160900 164731
		173850 207750 248600 258501
262	3q23	106165 110100 117700 150210 169600 180380 203500
		276902 601199 601682
265	19p13.1	143890 151440 600173 600276 600310 601604 601843
273	2p12-q22	104614 120435 126600 135300 136435 147200 152790
		157170 160980 178640 182601 203800 216900 253601
		278300 600678 601071 601771 602134 602404
274	15q22	102578 109700 151670 154550 601780
279	14q24.3	104311 109150 182600 245200 601208
281	1p33-p32	120260 120950 120960 138140 178300 187040 246450
		600101 600650 600722
283	1p36	118210 120550 120570 120575 121800 130500 133200
		155600 171760 185470 211420 230350 255800 601990
		602023 602771
287	9q21.32-q21.33	602014

288	2p23-p21	120435 126600 135300 136435 143450 152790 157170
200		182601 264600 278300 600890 601071 601771 602134
293	20	
301	6p21.3	106300 108800 120290 120810 120820 142857 142858
		150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
309	14	
312	1q32-q41	114208 119300 120620 120920 134370 134580 145260
		150310 179820 191045 276901 600105 600332 600759
315	Xq21.3-q22	601494 601744 601975 300088 300300 301201 301500 301835 303400 303630
313	Aq21.3-q22	303631 304500 304700 305450 309300 309605 311850
		312080
318	Xp22.32	306250 308100 312865
328	10p13	601362
332	17q21	109270 113705 144200 148065 148066 148067 148069
		148080 154275 168610 171190 176705 185800 200350
		221820 232200 249000 252920 253250 600119 601363
		601844
334	9q21.32-q21.33	602014
335	16q24	102600 103850 233690
337	11q22.3-q23	107680 107720 133780 147791 159555 168000 186740
,		186830 188025 203750 208900 261640 600048 601382
		602574
341	1p22-p31	170995 180069 191540 201450 248610 274270 600309
246	21-22.2	601414 601676 602094 602522
346 347	21q22.2 4p	176261 601399
348	12q23-q24	113100 124200 147440 158590 160781 163950 235800
340	12423-424	251170 276710 600175 601517
350	8q13-q22.1	122560 124080 170993 202010 214400 216550 222745
300	0415 422.1	259730 600415 601653 602476 602667
354	2p25	274500 602134
357	20q11.1-11.23	139190 224100 601002 601146
358	7q34-q35	118425 152427 180105 222800 274180 276000 600510
359	5q23-q31	121050 126150 131400 138040 153455 159000 179095
		181460 192974 600807 601596 601692 602089 602121
		602460
365	2cen-q24	
367	17q25	114290 138033 162100 170500 180860 264470
368	12q21.3-q22	147440 201470 235800 273300
371	14q24.3	104311 109150 182600 245200 601208
379	2q34	100730 118800 123660 135600 157655 186860 201460
501	20 11 2 15	205100 262000 601277 601318
381	20q11.2-q12	139190 224100 600281 601002 601146
384	7q22.1	120160 126650
386	17p13	138190 254210 271900 600179 600977 601202 601777
392	19p13.2	108725 120700 133171 143890 147670 151440 164953
		231670 600276 600957 601843

1210 134638 136132
9440 173610 176310
7400 233710 249270
1652 602491
0977 601202 601777
7000 212200 600624
7050 153700 161015
1181 193235 209901
0319 600528 601884
7670 151440 164953
0000 100071 101460
9000 180071 181460 2089
7700 600617
0820 142857 142858
9450 201910 217000
5550 600202 600261
000202 000201
4920 170650 600900
1920 170030 000300
8510 600919 601542
2280 600163 600971
2200 000103 0007/1
2221
4275 180240 182138
1954
2800 600965
1210 134638 136132
9440 173610 176310
7400 233710 249270
1652 602491
2005
2085
0011 (00701 (01600
0211 600701 601690
6490 171190 185800
0525 601844
9720 (01154 (01252
8720 601154 601253
6880 190195 222700
1369 602086 602279
8970
<del></del>
2760 314250 314580

566	11p15.5	125852 126452 141900 142000 142200 142250 142270
		176730 190020 191290 192500 194071 204500 600856
		601680 602631
567	9q13	190100 229300 600884 600974 602014
570	22q13.31	250100 250800
586	Xp11.23	300047 300071 300110 300600 301000 301830 309470
		309500 309610 309850 311050 312060
589	7p13-p12	138079 165240 180104 203740 219800 261670 601649
596	4q	
598	5q13.1	126060 143200 181510 600354
604	8q24.12-q24.13	133700 150230 190080 190350
606	7q36	142335 152427 163729 176450 190605 600510 600725
613	5q31-q32	109690 121050 131400 138040 138491 147061 147575
013	ا مامار	153455 154500 159000 179095 180071 181460 192974
		222600 272750 600807 601596 601692 602089 602121
		602460
632	Xq28	300031 300044 300048 300049 300055 300100 300104
	1-	300126 301201 301590 302060 302960 303700 303800
		303900 304800 305900 306700 306995 308310 308840
		309200 309548 309620 309900 310300 310400 310460
		311300 311510 314300 314400
635	6p21.3	106300 108800 120290 120810 120820 142857 142858
	1	150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
665	13q12.2	157900
676	19q13.2-q13.4	107741 113900 122720 126340 126391 130410 134790
		138570 152780 160900 164731 173850 191044 207750
		248600 258501 600040 600138 602225
682	19q13.1-q13.2	107741 113900 122720 126340 126391 160900 164731
	, ,	172400 173850 180901 207750 221770 248600 258501
		600918 602716
709	6q14-q21	120110 121014 136550 203310 269920 601666 602772
739	9p21	108120 112250 247640 600160 600221 601606
740	8q13-q21	122560 124080 202010 214400 600415 601653 602476
	1 - 1 -	602667
741	1p36.1-p35	116600 118210 120550 120570 120575 121800 130500
		133200 138140 138971 155600 171760 172411 185470
		211420 230350 255800 600975 601990 602023 602771
745	7q36	142335 152427 163729 176450 190605 600510 600725
746	6p22	248611
749	4q28-q32	107250 134820 134830 134850 181600 189800 208400
	11	231675 266300 600983
750	3p21.3	116806 120120 120436 138320 168468 182280 600163
767	3q23	106165 110100 117700 150210 169600 180380 203500
, 3,		276902 601199 601682
768	2q34	100730 118800 123660 135600 157655 186860 201460
, 00		205100 262000 601277 601318
776	2p24	107730 182601 602134
779	14q	
117	1 174	

PCT/US00/26524

780	9p11-p12	155900
781	Xq22	300088 300300 301201 301500 301835 303630 303631
		304500 304700 309300 309605 311850 312080
784	9q22	229600 264300 600542 602014 602088
788	5q23.3	121050 153455 159000 179095 192974 601596
790	1q32-q41	114208 119300 120620 120920 134370 134580 145260
		150310 179820 191045 276901 600105 600332 600759
		601494 601744 601975
797	5q31-q33	109690 121050 131400 138040 138491 147061 147575
		153455 154500 159000 179095 180071 181460 192974
		222600 234000 272750 600807 601411 601596 601692
		602089 602121 602460
814	21q22.3	120220 120240 123580 151385 171860 190685 236100
		236200 240300 267750 600065 601072 601145
815	12q13	107777 123940 139350 148040 148041 148043 148070
		231550 600194 600231 600536 600808 600956 601284
		601769 601928 602116 602153
828	12q24.2-q24.3	100650 142410 160781 181405
844	11p15.5	125852 126452 141900 142000 142200 142250 142270
		176730 190020 191290 192500 194071 204500 600856
		601680 602631
850	8q22	216550 259730
857	3p25	154705 193300 227646 253260 278720 601154 601253
		602011
881	1q41-q42	106150 145260 173870 276901 600332 600759 600996
		601744 601975
884	7q11.23	116860 129900 233700 600079
905	21q22.3	120220 120240 123580 151385 171860 190685 236100
		236200 240300 267750 600065 601072 601145
914	17p13.3	113721 247200 600059 601545
920	44	
933	17pter-p12	
951	4q31.3-q33	142380 189800 208400 231675
953	1q21	104770 107670 110700 135940 145001 146790 152445
		159001 174000 179755 182860 191315 230800 266200
		600897 601105 601412 601652 602491
957	20q12	600281
993	16q24.3	155555 227650 253000 602783
997	11p15	108985 186921 602092
1010	17q23	106180 138700 139250 150200 154275 176960 249000
		253250
1013	17p13.3	113721 247200 600059 601545
1031	20	
1048	11p15.5	125852 126452 141900 142000 142200 142250 142270
		176730 190020 191290 192500 194071 204500 600856
		601680 602631
1064	18q11.1-q11.2	114400 176300 257220 600192 600805
1070	2q21	133510 165320 223000
1071	5q23.3-q31.1	121050 131400 138040 147061 147575 153455 159000
	-	179095 181460 192974 600807 601596 601692 602089

		602121 602460
1072	12-14-2-15	181430 600698 600808 602116
1072	12q14.3-q15	
1074	17p13.3	113721 247200 600059 601545
1075	11	
1089	3p23-p21	116806 120120 120436 138320 139330 139360 150250
		164500 168468 182280 190182 212138 227646 230500
		238310 261510 600163 600971 601154 601226 601267
		601373 601916
1090	1p34.1	120550 120570 120575 121800 130500 133200 138140
		171760 178300 255800
1104	Xq22	300088 300300 301201 301500 301835 303630 303631
		304500 304700 309300 309605 311850 312080
1124	18q21	601941 602080 602397
1129	6q16.1-q16.3	136550 602772
1130	7q11.23	116860 129900 233700 600079
1131	11p15.4	130650 150000 257200
1153	19p13.3	108725 120700 133171 136836 145981 147141 164953
	1,512.0	188070 600957 601238 601846 602216 602477
1164	12p11-p13	103950 112410 113520 120580 131440 135700 138571
1104	12011 013	139130 142680 150100 168470 176260 186940 190070
		190450 200990 216950 600228 600414 600618 602096
1166	10p12	130 130 200330 210330 000220 000 111 000010 002030
1191	12q13	107777 123940 139350 148040 148041 148043 148070
1191	12413	231550 600194 600231 600536 600808 600956 601284
		601769 601928 602116 602153
1197	12p12.1-p11.2	112410 113520 135700 150100 168470 186940 190070
1197	12p12.1-p11.2	200990 602096
1198	1q21.2-q22	104770 107670 110700 145001 146760 146790 159440
1196	1421.2-422	186780 191030 191315 600923 601412 601652 601863
		602491
1199	2	002471
		102578 109700 118485 151670 154550 191010 231680
1214	15q22-q23	272800 276700 600374 601780
1240	1 21	
1240	1q21	104770 107670 110700 135940 145001 146790 152445 159001 174000 179755 182860 191315 230800 266200
1041	1 21	600897 601105 601412 601652 602491
1241	1q21	104770 107670 110700 135940 145001 146790 152445
		159001 174000 179755 182860 191315 230800 266200
1210	10.043	600897 601105 601412 601652 602491
1319	12q24.3	160781 181405
1325	Xp22.2	300075 300077 301200 302350 302801 305435 306000
		307800 308800 309510 311200 312040 312170 312700
		313400
1328	7	
1341	4	
1346	12q13-q15	107777 120140 123829 123940 126337 139350 147570
		148040 148041 148043 148070 181430 231550 232800
		252940 264700 600194 600231 600536 600698 600808
		600956 601284 601769 601928 602116 602153
1347	Xp22.2-p22.1	300029 300075 300077 301200 302350 302801 305435

		306000 306100 307800 308800 309510 311200 311770
		312040 312170 312700 313400
1357	15q14	102540 118511 218000 227220 243500 254770 601800
1360	6p21.3	106300 108800 120290 120810 120820 142857 142858
		150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
1397	19q13.2	107741 113900 122720 126340 126391 160900 164731
		173850 207750 248600 258501
1407	7p15-p14	107776 138079 139191 142959 153880 180104 203740
	7 15 14	600994 601472 601649
1418	7p15-p14	107776 138079 139191 142959 153880 180104 203740
1402	37 11 21	600994 601472 601649
1423	Xp11.21	300047 301300 301830 305400 308300 309470 309500
1426	12-22 -24 1	309610 311050   113100 124200 147440 158590 160781 163950 181405
1426	12q23-q24.1	235800 251170 261600 276710 600175 601406 601517
		601620 601621
1427	7p15-p14	107776 138079 139191 142959 153880 180104 203740
1427	/p15-p1 <del>4</del>	600994 601472 601649
1428	13q32-q33	133530 156600 232000 256731 601295 602085
1430	13q13	157900 600631
1435	7q35	118425 152427 180105 276000 600510
1438	5q13	126060 143200 181510 253200 268800 600354
1466	7q33	180105 222800
1471	10q11-q12	133540 154545 164761 188550 600835
1483	5q35.3	133340 134343 104701 188330 000833
1489	16	
1495	1q21.2-q21.3	104770 107670 110700 145001 146760 146790 191315
1473	1421.2-421.3	
1500	14a21-a22	The state of the s
		The state of the s
		the state of the s
1507	5q15.2-q21	
1509	14032.1	
1332	1921	
1536	6	
		108985 186921 602092
10.0	-4	
1542	1g32-g41	114208 119300 120620 120920 134370 134580 145260
	11	150310 179820 191045 276901 600105 600332 600759
		601494 601744 601975
1543	9p22	112250 147660 238300 247640
1545	15q21.3	109700 151670 601780
1547		129490 167415 176860 256100
1549		116600 118210 120550 120570 120575 121800 130500
1545 1547	11p15 11p15 1q21.1 1q32-q41	150310 179820 191045 276901 600105 600332 600759 601494 601744 601975  112250 147660 238300 247640  109700 151670 601780  129490 167415 176860 256100

-		133200 138140 138971 155600 171760 172411 185470
		211420 230350 255800 600975 601990 602023 602771
1550	1	
1556	11	
1562	17q24-q25	114290 115660 138033 139250 148500 150200 154275
		162100 170500 180860 182452 230200 249000 253250
		264470
1563	1q31	134580 145001 145260 150292 208250 226450 600105
		600759 600995 601652
1566	16p13.3	141750 141800 141850 156850 186580 191092 600140
		600273 601313 601785
1572	9q34.3	120215 190198
1574	2p15-cen	
1576	17q21-q22	109270 113705 120150 139250 144200 148065 148066
1		148067 148069 148080 150200 154275 156490 157140
		168610 171190 173470 176705 176960 185800 200350
		221820 232200 249000 252920 253250 273800 600119
		600525 600852 601363 601844
1579	21q11.2	159595
1580	21q11.2	159595
1581	22	
1582	12p13	103950 120580 131440 139130 142680 176260 190450
		200990 216950 600228 600414 600618 602096
1584	4p16	225500 600593 602363
1585	20	
1587	22q13.1	103050 124030 138981 182380 188826 190040
1590	18q11.2	114400 176300 257220 600192 600805
1591	7p15.3	153880 601649
1595	3p24	227646 600163 601154
1596	4q13-q21	103600 104150 104500 125490 147790 170650 173910
	• •	252500
1597	19p13.1	143890 151440 600173 600276 600310 601604 601843
1598	21q22.3	120220 120240 123580 151385 171860 190685 236100
	-	236200 240300 267750 600065 601072 601145
1600	20	
1603	12q13	107777 123940 139350 148040 148041 148043 148070
	•	231550 600194 600231 600536 600808 600956 601284
		601769 601928 602116 602153
1604	Xq28	300031 300044 300048 300049 300055 300100 300104
		300126 301201 301590 302060 302960 303700 303800
		303900 304800 305900 306700 306995 308310 308840
ļ.		309200 309548 309620 309900 310300 310400 310460
		311300 311510 314300 314400
1606	3q21-q22	106165 110100 117700 150210 169600 180380 190000
		203500 232050 276902 600882 601199 601471 601682
1609	8q21	124080 202010 214400 602476 602667
1613	9q31	109400 132800 186855 223900 253800 278700 602088
1614	11p15.5	125852 126452 141900 142000 142200 142250 142270
	-	176730 190020 191290 192500 194071 204500 600856
		601680 602631

1616		152200 167000 600200 600002 602544
1616	6q27	152200 167000 600320 600883 602544
1617	6q27	152200 167000 600320 600883 602544
1618	3	
1621	7p11	
1622	1q43	106150 156570 600996 601975 602759
1623	15q21-q22	102578 105600 107910 109700 114240 134797 151670
		154550 160777 600839 601780 602099
1626	20pter-p12	
1629	17	
1631	9q34.1	103000 114350 120900 131195 185000 189980 600184
		602575
1634	7p15	138079 139191 142959 153880 180104 600994 601649
1635	14q32.3	
1636	6q21	120110 121014 601666 602772
1637	Xq28	300031 300044 300048 300049 300055 300100 300104
		300126 301201 301590 302060 302960 303700 303800
		303900 304800 305900 306700 306995 308310 308840
		309200 309548 309620 309900 310300 310400 310460
		311300 311510 314300 314400
1645	13q12	121011 129500 253700 601885 602221
1646	12q22	147440 201470 235800 273300
1648	16q22	103850 114835 121360 217800 218030
1650	12	
1653	5q23	121050 126150 159000 179095 192974 601596
1656	14q24.1	182600
1660	3p21.1	150250 164500 168468 182280 238310 600163 601226
	•	601916
1664	14	
1665	3q13.3	126451 600882
1668	13q12.3-q13.1	157900 600185 600631
1669	13q12.3-q13.1	157900 600185 600631
1682	5q14.3	
1685		
	10q26	176943 258870 263700 601969 602084
1686		176943 258870 263700 601969 602084 150240 164160 180105 190900 222800 246900
1686 1691	7q31.3-q32	
1691	7q31.3-q32 7q31.2	150240 164160 180105 190900 222800 246900 150240 180105 222800 246900 602421
	7q31.3-q32 7q31.2 10q25	150240 164160 180105 190900 222800 246900
1691 1694	7q31.3-q32 7q31.2	150240 164160 180105 190900 222800 246900 150240 180105 222800 246900 602421 167409 278000 600020 600095 602669
1691 1694	7q31.3-q32 7q31.2 10q25	150240 164160 180105 190900 222800 246900 150240 180105 222800 246900 602421 167409 278000 600020 600095 602669 106300 108800 120290 120810 120820 142857 142858
1691 1694	7q31.3-q32 7q31.2 10q25	150240     164160     180105     190900     222800     246900       150240     180105     222800     246900     602421       167409     278000     600020     600095     602669       106300     108800     120290     120810     120820     142857     142858       150270     167250     170261     177900     179450     201910     217000
1691 1694	7q31.3-q32 7q31.2 10q25 6p21.3	150240     164160     180105     190900     222800     246900       150240     180105     222800     246900     602421       167409     278000     600020     600095     602669       106300     108800     120290     120810     120820     142857     142858       150270     167250     170261     177900     179450     201910     217000       222100     233100     235200     248611     256550     600202     600261
1691 1694 1702	7q31.3-q32 7q31.2 10q25 6p21.3	150240       164160       180105       190900       222800       246900         150240       180105       222800       246900       602421         167409       278000       600020       600095       602669         106300       108800       120290       120810       120820       142857       142858         150270       167250       170261       177900       179450       201910       217000         222100       233100       235200       248611       256550       600202       600261         601868       602280       602475
1691 1694 1702 1703 1708	7q31.3-q32 7q31.2 10q25 6p21.3	150240       164160       180105       190900       222800       246900         150240       180105       222800       246900       602421         167409       278000       600020       600095       602669         106300       108800       120290       120810       120820       142857       142858         150270       167250       170261       177900       179450       201910       217000         222100       233100       235200       248611       256550       600202       600261         601868       602280       602475         176000       261640       602574
1691 1694 1702 1703 1708 1710	7q31.3-q32 7q31.2 10q25 6p21.3 11q23.3 20p12 8q24-qter	150240       164160       180105       190900       222800       246900         150240       180105       222800       246900       602421         167409       278000       600020       600095       602669         106300       108800       120290       120810       120820       142857       142858         150270       167250       170261       177900       179450       201910       217000         222100       233100       235200       248611       256550       600202       600261         601868       602280       602475         176000       261640       602574         112261       176640       236700       601920
1691 1694 1702 1703 1708	7q31.3-q32 7q31.2 10q25 6p21.3	150240       164160       180105       190900       222800       246900         150240       180105       222800       246900       602421         167409       278000       600020       600095       602669         106300       108800       120290       120810       120820       142857       142858         150270       167250       170261       177900       179450       201910       217000         222100       233100       235200       248611       256550       600202       600261         601868       602280       602475         176000       261640       602574         112261       176640       236700       601920         125270       128100       137350       191100       215700       223360       268900
1691 1694 1702 1703 1708 1710 1711	7q31.3-q32 7q31.2 10q25 6p21.3 11q23.3 20p12 8q24-qter 9q34	150240       164160       180105       190900       222800       246900         150240       180105       222800       246900       602421         167409       278000       600020       600095       602669         106300       108800       120290       120810       120820       142857       142858         150270       167250       170261       177900       179450       201910       217000         222100       233100       235200       248611       256550       600202       600261         601868       602280       602475       601920         112261       176640       236700       601920         125270       128100       137350       191100       215700       223360       268900         601850
1691 1694 1702 1703 1708 1710	7q31.3-q32 7q31.2 10q25 6p21.3 11q23.3 20p12 8q24-qter	150240       164160       180105       190900       222800       246900         150240       180105       222800       246900       602421         167409       278000       600020       600095       602669         106300       108800       120290       120810       120820       142857       142858         150270       167250       170261       177900       179450       201910       217000         222100       233100       235200       248611       256550       600202       600261         601868       602280       602475       176000       261640       602574       112261       176640       236700       601920         125270       128100       137350       191100       215700       223360       268900         601850       141750       141800       141850       156850       186580       191092       600140
1691 1694 1702 1703 1708 1710 1711	7q31.3-q32 7q31.2 10q25 6p21.3  11q23.3 20p12 8q24-qter 9q34  16p13.3	150240       164160       180105       190900       222800       246900         150240       180105       222800       246900       602421         167409       278000       600020       600095       602669         106300       108800       120290       120810       120820       142857       142858         150270       167250       170261       177900       179450       201910       217000         222100       233100       235200       248611       256550       600202       600261         601868       602280       602475       601920         112261       176640       236700       601920         125270       128100       137350       191100       215700       223360       268900         601850
1691 1694 1702 1703 1708 1710 1711	7q31.3-q32 7q31.2 10q25 6p21.3 11q23.3 20p12 8q24-qter 9q34	150240       164160       180105       190900       222800       246900         150240       180105       222800       246900       602421         167409       278000       600020       600095       602669         106300       108800       120290       120810       120820       142857       142858         150270       167250       170261       177900       179450       201910       217000         222100       233100       235200       248611       256550       600202       600261         601868       602280       602475       176000       261640       602574       112261       176640       236700       601920         125270       128100       137350       191100       215700       223360       268900         601850       141750       141800       141850       156850       186580       191092       600140

1.505		T40650 4000 40000 04600 074600 600001
1793	7q3 <u>1</u>	126650 164860 180105 222800 246900 274600 602081
1794	7q31	126650 164860 180105 222800 246900 274600 602081
1804	11q13.1	106100 133780 601650
1811	2q11.1-q22	129490 133510 165320 167415 176860 176947 223000
		256030 256100
1818	9q22.3-q31	109400 132800 162400 186855 223900 227645 229700
	-	253800 278700 601309 602088
1834	7q21.3	129900 154276 173360 183600 602136 602447
1835	7q36	142335 152427 163729 176450 190605 600510 600725
1843	12p12	112410 113520 135700 168470 186940 200990 602096
1851	17q21.1	109270 148065 148080 154275 157140 171190 185800
	•	221820 249000 253250 600119 601844
1852	11p15	108985 186921 602092
1856	3q21-q25	106165 110100 117700 150210 169600 180380 190000
		203500 222900 232050 276902 600882 601199 601471
		601682
1864	21q22.3	120220 120240 123580 151385 171860 190685 236100
	<b>_</b>	236200 240300 267750 600065 601072 601145
1879	9q13	190100 229300 600884 600974 602014
1883	17p12-p11.2	100710 182290 201475 231200 262850 270200 600977
1002	7. P P	601097 601777 602666
1884	22q11.1-q11.2	123620 137181 138720 145410 231950 239500 275350
100.		600850
1890	11q22.3-q23.3	107680 107720 133780 147791 159555 168000 176000
1070	11422.5 425.5	176010 176797 186740 186830 188025 203750 208900
		261640 600048 601382 602574
1891	2p21-p16	104614 120435 126600 135300 136435 152790 157170
707	-P P	160980 182601 600678 601771
1893	3q13.1-q13.2	600467 600882
1894	7p13-p12	138079 165240 180104 203740 219800 261670 601649
1895	1	
1896	14q22	112262 182600 182870 232700 602086
1897	14q22	112262 182600 182870 232700 602086
1901	17p13.1-p11.2	100710 138190 182290 191170 201475 231200 254210
1,01	1, p. 5.1 p. 1.2	262850 270200 271900 600179 600977 601097 601202
		601777 602666
1906	2q14-q32	100690 120180 120190 133510 138430 142989 156232
.,,,,,	241, 402	165320 167415 176860 178600 223000 256030 266100
		278250 600258 600321 601847
1910	6p21.3	106300 108800 120290 120810 120820 142857 142858
	F	150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
1911	6p12	180297 230450 263200 601690
1916	5p14-p13	108962 120940 217050 217070 245050 600837 600946
1935	11q13	102200 106100 131100 133780 147050 153700 161015
	71	164009 168461 180721 180840 191181 193235 209901
		232600 259700 259770 600045 600319 600528 601884
1940	1	
1944	22	
		<u> </u>

1945	14	
1961	lp22-p31	170995 180069 191540 201450 248610 274270 600309
	r	601414 601676 602094 602522
1971	1q21.2-q22	104770 107670 110700 145001 146760 146790 159440
	- 4	186780 191030 191315 600923 601412 601652 601863
		602491
1973	16q13-q21	114835 118470 132700 172490 209900 600968
1975	5q35.3	
1979	10q24	157640 174900 180250 186770 236730 271245 278000
	•	600095 600512 601107 601130 602082
1983	15q	
1984	6p12	180297 230450 263200 601690
1986	17p13	138190 254210 271900 600179 600977 601202 601777
1989	12q24.31	181405
1993	6p21.3	106300 108800 120290 120810 120820 142857 142858
	•	150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
1994	22q11.23	123620 600850
1995	1p33	120260 138140 178300 246450
1996	11q22.3	133780 203750 208900 261640 602574
2008	17	
2011	12q24.2	100650 142410 160781 181405
2015	1q32	114208 119300 120620 120920 134370 134580 145260
	- 4	150310 179820 191045 600105 600759 601494 601975
2018	Xq22	300088 300300 301201 301500 301835 303630 303631
		304500 304700 309300 309605 311850 312080
2020	22q13.2-q13.31	188826 250100 250800
2021	5q33-q34	109690 123101 131400 154500 164770 180071 181460
		222600 234000 272750 600584 600807 601411 601596
		602089
2023	3p21.3	116806 120120 120436 138320 168468 182280 600163
2024	5q14-q21	143200 159350 162150 175100
2030	11p15.5	125852 126452 141900 142000 142200 142250 142270
		176730 190020 191290 192500 194071 204500 600856
		601680 602631
2042	11p14-p15.2	108985 114130 114550 168450 186921 257200 276904
20.2	7.p p.s.2	600509 600937 601387 602092
2043	17	
	- ·	
	19a13.4	134790 191044 600040 600138
2061	19q13.4 9a33-a34	134790 191044 600040 600138 125270 128100 137350 146150 191100 215700 223360
	19q13.4 9q33-q34	125270 128100 137350 146150 191100 215700 223360
2061 2064	9q33-q34	125270 128100 137350 146150 191100 215700 223360 223900 253800 268900 601850
2061 2064 2065	9q33-q34 15q15	125270 128100 137350 146150 191100 215700 223360 223900 253800 268900 601850 177070 182500 218000 227220 243500 600839 601800
2061 2064	9q33-q34	125270 128100 137350 146150 191100 215700 223360 223900 253800 268900 601850 177070 182500 218000 227220 243500 600839 601800 300031 300044 300048 300049 300055 300100 300104
2061 2064 2065	9q33-q34 15q15	125270 128100 137350 146150 191100 215700 223360 223900 253800 268900 601850 177070 182500 218000 227220 243500 600839 601800 300031 300044 300048 300049 300055 300100 300104 300126 301201 301590 302060 302960 303700 303800
2061 2064 2065	9q33-q34 15q15	125270 128100 137350 146150 191100 215700 223360 223900 253800 268900 601850 177070 182500 218000 227220 243500 600839 601800 300031 300044 300048 300049 300055 300100 300104 300126 301201 301590 302060 302960 303700 303800 303900 304800 305900 306700 306995 308310 308840
2061 2064 2065	9q33-q34 15q15	125270 128100 137350 146150 191100 215700 223360 223900 253800 268900 601850    177070 182500 218000 227220 243500 600839 601800   300031 300044 300048 300049 300055 300100 300104 300126 301201 301590 302060 302960 303700 303800 303900 304800 305900 306700 306995 308310 308840 309200 309548 309620 309900 310300 310400 310460
2061 2064 2065	9q33-q34 15q15	125270 128100 137350 146150 191100 215700 223360 223900 253800 268900 601850 177070 182500 218000 227220 243500 600839 601800 300031 300044 300048 300049 300055 300100 300104 300126 301201 301590 302060 302960 303700 303800 303900 304800 305900 306700 306995 308310 308840

2077	17	
2079	21q22.3	120220 120240 123580 151385 171860 190685 236100
		236200 240300 267750 600065 601072 601145
2080	17q21.3-q22	109270 120150 139250 148065 148080 150200 154275
1		156490 171190 173470 176960 185800 221820 249000
		253250 273800 600119 600525 600852 601844
2081	1q32	114208 119300 120620 120920 134370 134580 145260
	•	150310 179820 191045 600105 600759 601494 601975
2097	19q13.3	113900 126340 126391 130410 134790 138570 160900
		173850 258501 600040 602225
2098	2p15	
2100	11q24	600359 602574
2113	11q21-q22	105580 133780 203100 602574
2115	8p12-p11.2	136350 152760 173370 180100 182900 185430 270800
		277700 600617
2121	6p21.3	106300 108800 120290 120810 120820 142857 142858
		150270 167250 170261 177900 179450 201910 217000
ŀ		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
2125	12p12	112410 113520 135700 168470 186940 200990 602096
2129	2p24	107730 182601 602134
2130	11p11.2	133701 168500 171650 176930 600623 600811 600958
2131	Xq13.1	304040 305100 305450 309605 312760 314250 314580
2145	15q21-q22.2	102578 105600 107910 109700 114240 134797 151670
		154550 160777 191010 600839 601780 602099
2149	4p	
2153	12q	
2158	16q22	103850 114835 121360 217800 218030
2163	22q12.2	101000 123620 138981 188826 600850 601669
2180	19q12-q13.1	109560 164731 172400 180901 205900 221770 248600
		600652 600757 600918 602716
2181	11q22-q23	105580 107680 107720 133780 147791 159555 168000
		186740 186830 188025 203750 208900 261640 600048
	2 22 22	601382 602574
2183	2q22-q23	256030
2187	1q23	104770 107300 107670 131210 134638 136132 145001
		146740 146790 173610 176310 186780 191030 227400
2100	11012 2	601412 601652 602491 133780 151400
2188	11q13.3 12	133700 131400
2189		126550 202210 260020 602772
2193	6q14	136550 203310 269920 602772 160781 181405
2197	12q24.3	214400 600415 601653 602476
2199	8q13.3	116860 126650 129900 133170 154276 171050 171060
2200	7q13-q22	173360 183600 253220 602136 602447
2201	1.25	107300 131210 136132 145001 150292 173610 208250
2201	1q25	233710 600995 601518 601652
2203	22q11.2	123620 138720 145410 231950 239500 275350 600850
2203	11q24	600359 602574
2204	20q13.1	256540 600281
2203	20413.1	230340 000281

2207	15q13	103581 146150 218000 227220 601623 601800 601889 602117
2212	1p36	118210 120550 120570 120575 121800 130500 133200
2212	1000	155600 171760 185470 211420 230350 255800 601990
		602023 602771
2215	12q13-q15	107777 120140 123829 123940 126337 139350 147570
2213	12415-415	148040 148041 148043 148070 181430 231550 232800
		252940 264700 600194 600231 600536 600698 600808
		600956 601284 601769 601928 602116 602153
2217	11p13-p15	102772 106210 107271 108985 114550 115500 136530
	11p15 p15	151390 179615 179616 180385 186921 194070 245349
		602092
2227	21	
2232	12q23	124200 147440 160781 235800 600175
2239	16p13.3	141750 141800 141850 156850 186580 191092 600140
	-	600273 601313 601785
2240	20p11.2	121700 122000 188040
2266	2p12	147200 178640 216900
2267	2p12	147200 178640 216900
2268	2p22-p21	120435 126600 135300 136435 152790 157170 182601
		278300 601071 601771 602134
2270	22q13.33	
2271	22q13.31	250100 250800
2272	22q13.1	103050 124030 138981 182380 188826 190040
2280	15q22	102578 109700 151670 154550 601780
2281	5p14-p13	108962 120940 217050 217070 245050 600837 600946
2283	17q21	109270 113705 144200 148065 148066 148067 148069
	•	148080 154275 168610 171190 176705 185800 200350
		221820 232200 249000 252920 253250 600119 601363
		601844
2284	17q21	109270 113705 144200 148065 148066 148067 148069
	-	148080 154275 168610 171190 176705 185800 200350
		221820 232200 249000 252920 253250 600119 601363
		601844
2285	12q12-q14	107777 120140 123829 123940 126337 139350 147570
		148040 148041 148043 148070 181430 231550 232800
		252940 264700 600194 600231 600536 600808 600956
		601284 601769 601928 602116 602153
2286	16q24.3	155555 227650 253000 602783
2287	9q33-q34	125270 128100 137350 146150 191100 215700 223360
		223900 253800 268900 601850
2288	12	
2290	16q22	103850 114835 121360 217800 218030
2291	13q <sup>34</sup>	153900 180381 227500 227600 238970
2292	2q31	100690 120180 120190 142989 156232 178600 266100
		600258 600321
2293	1p34	130500 133200 138140 168360 171760 176100 178300
		230000 255800
2294	22q13.2-q13.31	188826 250100 250800
2298	11q13	102200 106100 131100 133780 147050 153700 161015

		164000 460461 100701 100040 101101 100006 000001
		164009 168461 180721 180840 191181 193235 209901
- 2200	11.15	232600 259700 259770 600045 600319 600528 601884
2299	11p15	108985 186921 602092
2300	3p21.1	150250 164500 168468 182280 238310 600163 601226
	11 02 03	601916
2303	11q22-q23	105580 107680 107720 133780 147791 159555 168000
		186740 186830 188025 203750 208900 261640 600048
2204		601382 602574
2304	4p	112721 247200 (00050 (01545
2305	17p13.3	113721 247200 600059 601545
2306	17p13.3	113721 247200 600059 601545
2307	10	110501 01500 (00050 (01515
2308	17p13.3	113721 247200 600059 601545
2310	15q21-q22.2	102578 105600 107910 109700 114240 134797 151670
	- 10	154550 160777 191010 600839 601780 602099
2312	11q13	102200 106100 131100 133780 147050 153700 161015
		164009 168461 180721 180840 191181 193235 209901
2202	0 11	232600 259700 259770 600045 600319 600528 601884
2392	8q11	600899
2403	8q21	124080 202010 214400 602476 602667
2404	19q13.1-q13.2	107741 113900 122720 126340 126391 160900 164731
		172400 173850 180901 207750 221770 248600 258501
2405		600918 602716
2405	6p	10.4770 107.770 110700 125040 145001 14.7700 152445
2408	1q21	104770 107670 110700 135940 145001 146790 152445
		159001 174000 179755 182860 191315 230800 266200
2416	1.1	600897 601105 601412 601652 602491
2416	11 12 2	112721 247200 (00050 (01545
2419	17p13.3	113721 247200 600059 601545
2420	17p13.3	113721 247200 600059 601545
2421	17p13.3	113721 247200 600059 601545
2422	17p13.3	113721 247200 600059 601545
2424	1p34.1-p32	120260 120550 120570 120575 120950 120960 121800
		130500 133200 138140 168360 171760 176100 178300
	1 241 22	187040 230000 246450 255800 600101 600650 600722
2425	lp34.1-p32	120260 120550 120570 120575 120950 120960 121800
		130500 133200 138140 168360 171760 176100 178300 187040 230000 246450 255800 600101 600650 600722
2426	1 241 .22	120260 120550 120570 120575 120950 120960 121800
2426	1p34.1-p32	
		130500 133200 138140 168360 171760 176100 178300
2427	1 24 1 22	187040 230000 246450 255800 600101 600650 600722
2427	1p34.1-p32	120260 120550 120570 120575 120950 120960 121800
		130500 133200 138140 168360 171760 176100 178300
2420	7-11 3	187040 230000 246450 255800 600101 600650 600722
2429	7q11.2	116860 129900 130160 173510
2431	12q13	107777 123940 139350 148040 148041 148043 148070
		231550 600194 600231 600536 600808 600956 601284
0.400	0.01	601769 601928 602116 602153
2433	8p21	152760 180100 185430 602629
2434	1q21	104770 107670 110700 135940 145001 146790 152445

		159001 174000 179755 182860 191315 230800 266200
	<del></del>	600897 601105 601412 601652 602491
2435	11p15	108985 186921 602092
2437	12q13.3	181430 232800 600808 601284 601769 602116
2438	4p16.3	134934 143100 180072 194190 252800 600965
2439	6p21.3	106300 108800 120290 120810 120820 142857 142858
		150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
2440	6q22-q23	107470 120110 121014 142470 156225 164200 207800
	16 12 2	601316 601410 601757 602067
2442	16p13.3	141750 141800 141850 156850 186580 191092 600140
	10	600273 601313 601785
2443	12	
2445	19pter-q12	102000 114250 120000 125270 120100 121105 127250
2446	9q33-q34.1	103000 114350 120900 125270 128100 131195 137350 146150 185000 189980 191100 215700 223360 223900
2447	2q12-q21	253800 268900 600184 601850 602575 129490 133510 165320 167415 176860 176947 223000
244 /	2q12-q21	256100
2448	1p34.1	120550 120570 120575 121800 130500 133200 138140
2448	1934.1	171760 178300 255800
2449	40	171700 178300 233800
2450	4q 7p	
2451		118425 152427 180105 276000 600510
	7q35	102200 106100 131100 133780 147050 153700 161015
2452	11q13	164009 168461 180721 180840 191181 193235 209901
		232600 259700 259770 600045 600319 600528 601884
2453	2q23	232000 239700 239770 000043 000319 000328 001004
2454	0cen a31	
2454	9cen-q34	182600
2456	14q24.1	182600
2456 2457	14q24.1 9q12	602014
2456 2457 2463	14q24.1 9q12 20q13.2-q13.3	602014 118504 131242 139320 602025 602235
2456 2457	14q24.1 9q12	602014 118504 131242 139320 602025 602235 104770 107670 110700 145001 146760 146790 159440
2456 2457 2463	14q24.1 9q12 20q13.2-q13.3	602014 118504 131242 139320 602025 602235 104770 107670 110700 145001 146760 146790 159440 186780 191030 191315 600923 601412 601652 601863
2456 2457 2463 2466	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22	602014 118504 131242 139320 602025 602235 104770 107670 110700 145001 146760 146790 159440 186780 191030 191315 600923 601412 601652 601863 602491
2456 2457 2463 2466	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22	602014 118504 131242 139320 602025 602235 104770 107670 110700 145001 146760 146790 159440 186780 191030 191315 600923 601412 601652 601863 602491 107250 181600 189800 266300
2456 2457 2463 2466	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22	602014 118504 131242 139320 602025 602235 104770 107670 110700 145001 146760 146790 159440 186780 191030 191315 600923 601412 601652 601863 602491 107250 181600 189800 266300 139330 139360 144700 150250 156845 164500 182280
2456 2457 2463 2466 2467 2468	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22 4q31 3p21-p14	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491         107250       181600       189800       266300         139330       139360       144700       150250       156845       164500       182280         277730       600163       600971       601226       601267       601373
2456 2457 2463 2466 2467 2468 2469	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22 4q31 3p21-p14 4q24	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491         107250       181600       189800       266300         139330       139360       144700       150250       156845       164500       182280         277730       600163       600971       601226       601267       601373         157147       248510
2456 2457 2463 2466 2467 2468 2469 2470	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22 4q31 3p21-p14 4q24 13q12	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491       602
2456 2457 2463 2466 2467 2468 2469 2470 2471	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22 4q31 3p21-p14 4q24 13q12 2q21	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491       602491       602491       602491       602491       602491       602491       602491       602201       602201       602201       602201       602201       602201       602201       602201       602201       602201       602201       602000       602
2456 2457 2463 2466 2467 2468 2469 2470 2471 2476	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22 4q31 3p21-p14 4q24 13q12 2q21 7q34	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491       107250       181600       189800       266300         139330       139360       144700       150250       156845       164500       182280         277730       600163       600971       601226       601267       601373         157147       248510         121011       129500       253700       601885       602221         133510       165320       223000         180105       222800       274180
2456 2457 2463 2466 2467 2468 2469 2470 2471 2476 2477	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22  4q31 3p21-p14  4q24 13q12 2q21 7q34 6q14	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491       602491       602491       602491       602491       602491       602491       602491       602201       602201       602201       602201       602201       602201       602201       602201       602201       602201       602201       602000       602
2456 2457 2463 2466 2467 2468 2469 2470 2471 2476 2477 2478	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22  4q31 3p21-p14  4q24 13q12 2q21 7q34 6q14 12q	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491       107250       181600       189800       266300         139330       139360       144700       150250       156845       164500       182280         277730       600163       600971       601226       601267       601373         157147       248510         121011       129500       253700       601885       602221         133510       165320       223000         180105       222800       274180
2456 2457 2463 2466 2467 2468 2469 2470 2471 2476 2477 2478 2480	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22  4q31 3p21-p14  4q24 13q12 2q21 7q34 6q14 12q 22q13.33	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491       602491       602491       602491       602491       602491       602491       602201       602201       602201       602201       602201       602201       602201       602201       602201       602201       602000       602
2456 2457 2463 2466 2467 2468 2469 2470 2471 2476 2477 2478 2480 2485	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22  4q31 3p21-p14  4q24 13q12 2q21 7q34 6q14 12q 22q13.33 2p12	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491       107250       181600       189800       266300         139330       139360       144700       150250       156845       164500       182280         277730       600163       600971       601226       601267       601373         157147       248510         121011       129500       253700       601885       602221         133510       165320       223000         180105       222800       274180         136550       203310       269920       602772
2456 2457 2463 2466 2467 2468 2469 2470 2471 2476 2477 2478 2480	14q24.1 9q12 20q13.2-q13.3 1q21.2-q22  4q31 3p21-p14  4q24 13q12 2q21 7q34 6q14 12q 22q13.33	602014         118504       131242       139320       602025       602235         104770       107670       110700       145001       146760       146790       159440         186780       191030       191315       600923       601412       601652       601863         602491       602491       602491       602491       602491       602491       602491       602201       602201       602201       602201       602201       602201       602201       602201       602201       602201       602000       602

2490	22q11.1-q11.2	123620 137181 138720 145410 231950 239500 275350 600850
2491	22q11.1-q11.2	123620 137181 138720 145410 231950 239500 275350 600850
2495	2p12	147200 178640 216900
2496	2p12	147200 178640 216900
2497	2p12	147200 178640 216900
2498	22q11.1-q11.2	123620 137181 138720 145410 231950 239500 275350
2470	22q11.1-q11.2	600850
2499	14q32.33	144120 147020 147110
2501	22q11.1-q11.2	123620 137181 138720 145410 231950 239500 275350
2301	22q11.1-q11.2	600850
2513	17p13-p12	100710 138190 231200 254210 262850 271900 600179
2313	1/p13-p12	600977 601202 601777
2514	2q33-q36	100730 118800 120070 120131 123660 125660 135600
2314	2433-430	138030 147545 157655 186860 193500 201460 205100
	: :	213700 237300 259900 262000 600258 600266 601277
		601318
2515	2q32.2	600258 602087
2516	14q11.2	182600 186880 190195 222700 600243 602279
		107250 181600 189800 266300
2518	4q31	
2521	14q11.2-q21	135750 160760 164050 182600 186880 190195 222700
		232700 600243 600635 600792 601369 602086 602279
2524	1.5	602616
2524	17	110200 10000 100121 120000 120020 050000 (00420
2526	2q37	113300 120070 120131 120250 138030 259900 600430
2529	12p12.2-p12.1	112410 138571 150100 168470 190070 200990 602096
2532	2p12	147200 178640 216900
2533	12q21.3-q22	147440 201470 235800 273300
2535	11p15.5	125852 126452 141900 142000 142200 142250 142270
		176730 190020 191290 192500 194071 204500 600856
		601680 602631
2536	11	
2538	12p13	103950 120580 131440 139130 142680 176260 190450
		200990 216950 600228 600414 600618 602096
2539	6p21.3	106300 108800 120290 120810 120820 142857 142858
		150270 167250 170261 177900 179450 201910 217000
		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
2540	17q25	114290 138033 162100 170500 180860 264470
2542	2	
2543	17q	
2544	13q12	121011 129500 253700 601885 602221
2545	13q12	121011 129500 253700 601885 602221
2547	5q31.1	131400 147061 147575 153455 159000 181460 600807
	_	601596 602089
2548	2p22-p21	120435 126600 135300 136435 152790 157170 182601
	'	278300 601071 601771 602134
2554	2q21	133510 165320 223000
2555	Xq28	300031 300044 300048 300049 300055 300100 300104
		I control to the cont

		300126 301201 301590 302060 302960 303700 303800
		303900 304800 305900 306700 306995 308310 308840
		309200 309548 309620 309900 310300 310400 310460
		311300 311510 314300 314400
2561	1q21.2-q22	104770 107670 110700 145001 146760 146790 159440
		186780 191030 191315 600923 601412 601652 601863
		602491
2562	1q21.2-q22	104770 107670 110700 145001 146760 146790 159440
		186780 191030 191315 600923 601412 601652 601863
		602491
2564	9q34	125270 128100 137350 191100 215700 223360 268900
		601850
2565	9q34	125270 128100 137350 191100 215700 223360 268900
		601850
2568	5q35	123101 164040 208100 246530
2569	2q12-qter	
2570	1p35	118210 120550 120570 120575 121800 130500 133200
	-	138140 138971 171760 172411 185470 230350 255800
		602771
2576	12q13	107777 123940 139350 148040 148041 148043 148070
		231550 600194 600231 600536 600808 600956 601284
		601769 601928 602116 602153
2577	10q21.1	129010 601386 601493
2578	7p13-p11.2	138079 165240 180104 203740 219800 261670 601649
2579	10q22	126090 129010 142600 250850 601386 601493
2581	12q23	124200 147440 160781 235800 600175
2582	12q13.3	181430 232800 600808 601284 601769 602116
2585	15q22	102578 109700 151670 154550 601780
2587	5p14-p12	108962 120940 217050 217070 245050 600837 600946
2588	1q31-q32	114208 119300 120620 120920 134370 134580 145001
2300	1451 452	145260 150292 150310 179820 191045 208250 226450
		600105 600759 600995 601494 601652 601975
2590	8p22	148370 238600 600143 601385 602629
2592	4q12	103600 104150 104500 164920 170650 600900
2594	8p12-p11.2	136350 152760 173370 180100 182900 185430 270800
<b>-</b> 37 (	Op12 p11.2	277700 600617
2595	14q22	112262 182600 182870 232700 602086
2598	15q26.1	166800 210900
2599	Xq28	300031 300044 300048 300049 300055 300100 300104
2377	23420	300126 301201 301590 302060 302960 303700 303800
		303900 304800 305900 306700 306995 308310 308840
		309200 309548 309620 309900 310300 310400 310460
		311300 311510 314300 314400
2601	11q13.1-q13.3	106100 133780 151400 601650 602078
2602	17q21.2	109270 148065 148080 154275 171190 185800 221820
2002	1/421.2	249000 253250 600119 601844
2603	7q36	142335 152427 163729 176450 190605 600510 600725
2605	17q	11233 132421 103123 110430 130003 000310 000723
2606	1p36.1-p35	116600 118210 120550 120570 120575 121800 130500
2000	1p50.1-p55	133200 138140 138971 155600 171760 172411 185470
		133400 136140 1367/1 133000 1/1/00 1/2411 1834/0

		211420 230350 255800 600975 601990 602023 602771
2607	13q	211420 230330 233800 000973 001990 002023 002771
2608	17q11	147557 600881 601954
2609	17	147557 000001 001754
2610	19q13.4	134790 191044 600040 600138
2611	2	134770 171044 000040 000130
2612	4	
2613	16p13.3	141750 141800 141850 156850 186580 191092 600140
2013	10013.3	600273 601313 601785
2614	12q	000273 001313 001703
2615	14q	
2616	9p21	108120 112250 247640 600160 600221 601606
2617	19q12-q13.1	109560 164731 172400 180901 205900 221770 248600
2017	19412-415.1	600652 600757 600918 602716
2619	22q13.1	103050 124030 138981 182380 188826 190040
2620	4p16	225500 600593 602363
2621	16p13.11	145505 186580 278760
2623	6p21.3	106300 108800 120290 120810 120820 142857 142858
2023	Op21.5	150270 167250 170261 177900 179450 201910 217000
	,	222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
2626	21q22.3	120220 120240 123580 151385 171860 190685 236100
2020	21422.3	236200 240300 267750 600065 601072 601145
2627	1q31-q32	114208 119300 120620 120920 134370 134580 145001
2027	1431-432	145260 150292 150310 179820 191045 208250 226450
		600105 600759 600995 601494 601652 601975
2628	11p15.5-p15.4	125852 126452 130650 141900 142000 142200 142250
2020	11p15.5-p15.4	142270 150000 176730 190020 191290 192500 194071
		204500 257200 600856 601680 602631
2629	1	
2633	8q22-q23	148900 216550 259730
2634	12q12	600194 600231 601284 601769
2635	2p22-p21	120435 126600 135300 136435 152790 157170 182601
2033	2p22 p21	278300 601071 601771 602134
2636	7q21.1	129900 154276 171050 171060 602136 602447
2637	17p13.3	113721 247200 600059 601545
2640	2p12	147200 178640 216900
2641	5q11.2-q13.1	126060 143200 181510 214300 253200 268800 600354
_0 . L	71.1.2 415.1	600887
2642	22q13.1	103050 124030 138981 182380 188826 190040
2645	11q13.2-q13.3	133780 151400 602078
2648	5p13-cen	155,05 151100 0020,0
2650	14	
2651	12q24.2	100650 142410 160781 181405
2652	12q24.2 12q13.1	126337 600808 601284 601769 602116
2655	13q14	109543 600631 601499
2656		192340 234200
2657	20p13	192340 234200 141750 141800 141850 156850 186580 191092 600140
2037	16p13.3	600273 601313 601785
2650	10n12 1 n12	143890 151440 600173 600276 600310 601011 601604
2659	19p13.1-p12	143670 131440 000173 000270 000310 001011 001004

- 1		601768 601843
2660	1q41-q42	106150 145260 173870 276901 600332 600759 600996
2000	1941-942	601744 601975
2681	8p22	148370 238600 600143 601385 602629
2682	19p13.3-p13.2	108725 120700 133171 136836 143890 145981 147141
2002	19p15.5-p15.2	147670 151440 164953 188070 231670 600276 600957
		601238 601843 601846 602216 602477
2742	6p21.3	106300 108800 120290 120810 120820 142857 142858
2/42	op21.5	150270 167250 170261 177900 179450 201910 217000
ŀ		222100 233100 235200 248611 256550 600202 600261
		601868 602280 602475
2744	22q12.1	123620 188826 600850 601669
2786	19q13.2-q13.4	107741 113900 122720 126340 126391 130410 134790
2700	19413.2-413.4	138570 152780 160900 164731 173850 191044 207750
		248600 258501 600040 600138 602225
2808	17p13-p12	100710 138190 231200 254210 262850 271900 600179
2000	17p15 p12	600977 601202 601777
2815	3q21-q23	106165 110100 117700 150210 169600 180380 190000
2013	5 <b>42</b> 1 <b>42</b> 5	203500 232050 276902 600882 601199 601471 601682
2844	11p13	102772 106210 107271 114550 115500 136530 151390
2011	11915	179615 179616 180385 194070 245349
2849	5q14-q22	143200 159350 162150 175100
2858	6p21.1-p12	179605 180297 230450 248611 251000 263200 600211
2030	op21.1-p12	600364 600701 601498 601690
2863	10q26.3	263700
2869	12	203700
2901	2p12-p13	147200 178640 203800 216900 602404
2922	X	11/200 1/00 10 203000 210300 002101
2930	7q34	180105 222800 274180
2936	6q14	136550 203310 269920 602772
2942	14q11.2	182600 186880 190195 222700 600243 602279
2979	12q13	107777 123940 139350 148040 148041 148043 148070
2,,,	.54.0	231550 600194 600231 600536 600808 600956 601284
		601769 601928 602116 602153
3024	12p13	103950 120580 131440 139130 142680 176260 190450
	<b>F</b>	200990 216950 600228 600414 600618 602096
3070	5q31.1	131400 147061 147575 153455 159000 181460 600807
	. 1	601596 602089
3079	6q14	136550 203310 269920 602772
3080	6q14	136550 203310 269920 602772
3082	7	
3117	1p36	118210 120550 120570 120575 121800 130500 133200
	Ľ	155600 171760 185470 211420 230350 255800 601990
		602023 602771
3130	9p21	108120 112250 247640 600160 600221 601606
3133	11q25	602782
3154	12q24.1	124200 147440 160781 181405 261600 601406 601620
'	<b>-1</b>	601621
· · · · ·	V-22.22	306250 308100 312865
3177	Xp22.32	300230 300100 312003

	<u></u>	
3183	11q13	102200 106100 131100 133780 147050 153700 161015
		164009 168461 180721 180840 191181 193235 209901
		232600 259700 259770 600045 600319 600528 601884
3187	13q14	109543 600631 601499
3205	17p13.3	113721 247200 600059 601545
3226	5q14-q22	143200 159350 162150 175100
3236	11q13	102200 106100 131100 133780 147050 153700 161015
	•	164009 168461 180721 180840 191181 193235 209901
		232600 259700 259770 600045 600319 600528 601884
3243	17q21	109270 113705 144200 148065 148066 148067 148069
	•	148080 154275 168610 171190 176705 185800 200350
		221820 232200 249000 252920 253250 600119 601363
		601844
3259	13q14	109543 600631 601499
3281	14q24-q31	104311 107970 109150 115650 182600 245200 275200
		601208 602091
3287	19q13.3	113900 126340 126391 130410 134790 138570 160900
	•	173850 258501 600040 602225
3299	12q13-q14	107777 120140 123829 123940 126337 139350 147570
		148040 148041 148043 148070 181430 231550 232800
		252940 264700 600194 600231 600536 600808 600956
		601284 601769 601928 602116 602153
3306	2p25	274500 602134
3316	2	
3323	22q13.2-q13.31	188826 250100 250800
3329	21q22.1	147450 176261 253270 601399
3335	15q15.3	114240 224120 600839 602099
3351	12q12-q13	107777 123940 139350 148040 148041 148043 148070
		231550 600194 600231 600536 600808 600956 601284
		601769 601928 602116 602153
3357	9q11-q22	190100 200150 229300 229600 264300 600429 600542
		600884 600974 600998 602014 602088
3382	16q22.1	103850 114835 116800 140100 192090 245900 276600
		600223
3392	19q13.4	134790 191044 600040 600138
3411	5q13	126060 143200 181510 253200 268800 600354
3429	9q22.3	162400 227645 229700 278700 601309 602088
3434	6p21	180297 248611 251000 263200 600211 600701 601690
3439	15q15	177070 182500 218000 227220 243500 600839 601800
3442	20q12	600281
3445	12	
3451	2q31	100690 120180 120190 142989 156232 178600 266100
	<b>^</b>	600258 600321
3455	16q22	103850 114835 121360 217800 218030
3460	11q12	106100 147050 259700 259770 600045 601884
3465	11q13-q14	102200 106100 131100 133780 147050 151400 153700
	1 - 1	161015 164009 168461 180721 180840 191181 193235
		203100 209901 232600 245000 259700 259770 266150
		276903 600045 600319 600528 601650 601884 602078
3477	1q12-1q21.2	104770 107670 110700 135940 145001 146760 146790

		152445 159001 174000 179755 182860 191315 230800
2402	16.00.1	266200 600897 601105 601412 601652 601863 602491
3492	16q22.1	103850 114835 116800 140100 192090 245900 276600 600223
3497	17q25	114290 138033 162100 170500 180860 264470
3503	1p31-p12	102770 120280 164790 166600 170995 180069 188540
	•	191540 201450 201810 232400 248610 274270 600234
		600309 601414 601676 601691 601718 602094 602522
3526	20	
3532	5q23-31	121050 126150 131400 138040 153455 159000 179095
		181460 192974 600807 601596 601692 602089 602121
		602460
3544	2p25.2-p25.1	100000000000000000000000000000000000000
3549	12q24	113100 124200 147440 158590 160781 163950 251170 276710 600175 601517
3551	12q13	107777 123940 139350 148040 148041 148043 148070
3331	12415	231550 600194 600231 600536 600808 600956 601284
		601769 601928 602116 602153
3555	1p31	180069 201450 248610 600309 601676 602522
3559	5q33-q34	109690 123101 131400 154500 164770 180071 181460
		222600 234000 272750 600584 600807 601411 601596
		602089
3560	3q13.1-q13.2	600467 600882
3561	19q12	
3564	21q22.3	120220 120240 123580 151385 171860 190685 236100
		236200 240300 267750 600065 601072 601145
3566	16q12-q13	114835 132700 172490 600968 602218 602639
3567	16q24.3	155555 227650 253000 602783
3573	1q32	114208 119300 120620 120920 134370 134580 145260
	10 01 1	150310 179820 191045 600105 600759 601494 601975
3574	10q21.1	129010 601386 601493
3576	19	113900 126340 126391 130410 134790 138570 152780
3600	19q13.3-q13.4	160900 173850 191044 258501 600040 600138 602225
3607	12q13	107777 123940 139350 148040 148041 148043 148070
3007	12415	231550 600194 600231 600536 600808 600956 601284
		601769 601928 602116 602153
3608	1p33-p34	120260 130500 133200 138140 168360 171760 176100
	-P F-	178300 230000 246450 255800
3611	2	
3623	17q25	114290 138033 162100 170500 180860 264470
3636	7q11	
3646	3p21.3	116806 120120 120436 138320 168468 182280 600163
3647	11q13	102200 106100 131100 133780 147050 153700 161015
		164009 168461 180721 180840 191181 193235 209901
		232600 259700 259770 600045 600319 600528 601884
3650	Xp22.1	300075 300077 301200 302350 306000 306100 307800
2652	5 00 00	309510 311770 312040 312170 312700 313400
3652	5q22-q23	121050 126150 159000 175100 179095 192974 601596
3653	1q44-qter	

2650	10-12 1	164721 172400 180001 221770 248600 600018 602716
3659	19q13.1	164731 172400 180901 221770 248600 600918 602716
3671	7p21-p15	138079 139191 142959 153880 180104 600994 601622
- 2602	15.06	601649
3683	15q26	180090 600318
3688	10q11.2	154545 164761 188550
3690	12q24.31	181405
3691	6q14	136550 203310 269920 602772
3701	15q21-q22.2	102578 105600 107910 109700 114240 134797 151670
		154550 160777 191010 600839 601780 602099
3702	1	200045 20045 201005 201045 205150 210400
3703	Xq24	300046 300123 301201 301835 301845 307150 310490
		311850
3704	8q21.3-q22.1	216550 222745 259730
3705	2q31	100690 120180 120190 142989 156232 178600 266100
		600258 600321
3706	22q13.31	250100 250800
3707	3q12-q13	121300 146200 190300 258900 600882
3711	12q22-q23	124200 147440 160781 201470 235800 273300 600175
3712	15q21-q22.2	102578 105600 107910 109700 114240 134797 151670
		154550 160777 191010 600839 601780 602099
3729	8q	
3749	19q13.1-q13.2	107741 113900 122720 126340 126391 160900 164731
		172400 173850 180901 207750 221770 248600 258501
		600918 602716
3773	12q24.2	100650 142410 160781 181405
3782	3p21	139330 139360 150250 164500 182280 600163 600971
		601226 601267 601373
3784	5q13.3 <u>-q</u> 14	139150 143200 181510 600354
3800	4q13-q21	103600 104150 104500 125490 147790 170650 173910
2802	C-14	252500 136550 203310 269920 602772
3803	6q14	165215 222900 600049
3831	3q26	
3838	10p11.2	600964 602026 102770 120280 166600 170995 232400 600309 601414
3854	1p21	
2062	10 01 1	601691 601718 602094 174810 600624 600993 602080
3863	18q21.1	1/4810 600624 600993 602080
3864	6q	100705 120700 122171 126926 142900 145091 147141
3871	19p13.3-p13.2	108725 120700 133171 136836 143890 145981 147141 147670 151440 164953 188070 231670 600276 600957
		601238 601843 601846 602216 602477
2077	2 21 2 21 1	
3877	2p21.3-p21.1	120435 182601 601771   107741 113900 122720 126340 126391 160900 164731
3879	19q13.1-q13.2	
		172400 173850 180901 207750 221770 248600 258501 600918 602716
3887	4-01 -05	103720 104500 125490 137600 138850 147790 157147
388/	4q21-q25	163890 173910 189800 217030 248510 252500 600919
		601542
3888	1q21-q23	104770 107300 107670 110700 131210 134638 135940
3000	1421-423	136132 145001 146740 146760 146790 152445 159001
		159440 173610 174000 176310 179755 182860 186780
		137770 173010 177000 170310 177733 102000 100700

		191030 191315 227400 230800 266200 600897 600923
		601105 601412 601652 601863 602491
3908	15q22.3-q23	118485 151670 231680 272800 276700 600374 601780
3911	1q42.1	106150 136850 214500 600996 601975
3917	13	
3918	17q24.3-q25.1	114290 138033 162100 170500 180860 264470
3919	11q14.1-q14.3	133780 203100
3923	1pter-p35	
3926	10q22	126090 129010 142600 250850 601386 601493
3930	17p13.3	113721 247200 600059 601545
3971	4q	113721 217200 000009 0012 10
3977	15q15	177070 182500 218000 227220 243500 600839 601800
3993	19p13.3	108725 120700 133171 136836 145981 147141 164953
3993	19015.5	188070 600957 601238 601846 602216 602477
4001	Xq26.1-q27.2	300085 300123 300700 301201 301590 301845 301900
4001	Aq20.1-q27.2	304340 306900 306955 307150 307700 308000 309000
		310490 313850
4002	17-12 2	113721 247200 600059 601545
4003	17p13.3	103950 120580 131440 139130 142680 176260 190450
4008	12p13	
4011	16.00.1	200990 216950 600228 600414 600618 602096
4011	16q22.1	103850 114835 116800 140100 192090 245900 276600
		600223
4013	10	
4018	16q22.1	103850 114835 116800 140100 192090 245900 276600 600223
4029	6q21-q22	120110 121014 156225 164200 601410 601666 601757
,,,,,	V-1	602772
4047	1p32-p31	120950 120960 138140 178300 180069 187040 201450
	-	248610 600101 600309 600650 600722 601676 602522
4054	11q23	107680 107720 133780 147791 159555 168000 186740
	·	186830 188025 203750 261640 600048 601382 602574
4058	12q22-qter	
4061	5q31.3-q32	109690 131400 138491 154500 159000 180071 181460
	• •	222600 272750 600807 601596 602089
4085	16q22.1	103850 114835 116800 140100 192090 245900 276600
	•	600223
4093	19q13.4	134790 191044 600040 600138
4100	10q25	167409 278000 600020 600095 602669
4105	17q21	109270 113705 144200 148065 148066 148067 148069
	- , 4	148080 154275 168610 171190 176705 185800 200350
		221820 232200 249000 252920 253250 600119 601363
		601844
4125	4q21	104500 125490 147790 173910 252500
4128	17p13	138190 254210 271900 600179 600977 601202 601777
4143	19q13.2	107741 113900 122720 126340 126391 160900 164731
ברוד <i>י</i>	17413.2	173850 207750 248600 258501
4149	4q27	147680 189800 600919
	11p15.4	130650 150000 257200
4171		
4178	12q22-q23	124200 147440 160781 201470 235800 273300 600175
4185	13q33	133530 601295

4192	1q31	134580 145001 145260 150292 208250 226450 600105
		600759 600995 601652
4196	8q24.3	188450
4245	2p12	147200 178640 216900
4261	12p13	103950 120580 131440 139130 142680 176260 190450
		200990 216950 600228 600414 600618 602096
4262	12q13-q15	107777 120140 123829 123940 126337 139350 147570
		148040 148041 148043 148070 181430 231550 232800
		252940 264700 600194 600231 600536 600698 600808
		600956 601284 601769 601928 602116 602153
4263	11q22.2-q22.3	133780 203750 208900 261640 602574
4270	1q21.2-q22	104770 107670 110700 145001 146760 146790 159440
		186780 191030 191315 600923 601412 601652 601863
		602491

Table 5

Library	Library Description
Code	Morton Fetal
H0002	Human Adult Heart
H0004	Human Adult Spleen
H0007	Human Cerebellum
H0008	Whole 6 Week Old Embryo
H0009	Human Fetal Brain
H0011	Human Fetal Kidney
H0011	Human Fetal Kidney
H0012	Human 8 Week Whole Embryo
H0013	Human Gall Bladder
H0015	Human Gall Bladder, fraction II
H0019	Human Fetal Heart
H0022	Jurkat Cells
H0023	Human fetal lung
H0024	Human Fetal Lung III
H0024	Namalwa Cells
H0030	Human Placenta
H0031	Human Placenta
H0031	Human Prostate
H0036	Human Adult Small Intestine
H0037	Human Adult Small Intestine
H0037	Human Testes
H0039	Human Pancreas Tumor
H0040	Human Testes Tumor
H0041	Human Fetal Bone
H0042	Human Adult Pulmonary
H0044	Human Cornea
H0045	Human Esophagus, Cancer
H0046	Human Endometrial Tumor
H0048	Human Pineal Gland
H0050	Human Fetal Heart
H0051	Human Hippocampus
H0052	Human Cerebellum
H0056	Human Umbilical Vein, Endo. remake
H0057	Human Fetal Spleen
H0059	Human Uterine Cancer
H0063	Human Thymus
H0068	Human Skin Tumor
H0069	Human Activated T-Cells
H0071	Human Infant Adrenal Gland
H0074	Human Platelets
H0081	Human Fetal Epithelium (Skin)
H0083	HUMAN JURKAT MEMBRANE BOUND POLYSOMES
H0085	Human Colon
H0085	Human epithelioid sarcoma

H0087	Human Thymus
H0090	Human T-Cell Lymphoma
H0097	Human Adult Heart, subtracted
H0098	Human Adult Liver, subtracted
H0100	Human Whole Six Week Old Embryo
H0101	Human 7 Weeks Old Embryo, subtracted
H0102	Human Whole 6 Week Old Embryo (II), subt
H0105	Human Fetal Heart, subtracted
H0107	Human Infant Adrenal Gland, subtracted
H0108	Human Adult Lymph Node, subtracted
H0116	Human Thymus Tumor, subtracted
H0119	Human Pediatric Kidney
H0122	Human Adult Skeletal Muscle
H0123	Human Fetal Dura Mater
H0124	Human Rhabdomyosarcoma
H0125	Cem cells cyclohexamide treated
H0130	LNCAP untreated
H0131	LNCAP + o.3nM R1881
H0132	LNCAP + 30nM R1881
H0134	Raji Cells, cyclohexamide treated
H0135	Human Synovial Sarcoma
H0136	Supt Cells, cyclohexamide treated
H0144	Nine Week Old Early Stage Human
H0147	Human Adult Liver
H0150	Human Epididymus
H0153	Human adult lymph node, subtracted
H0156	Human Adrenal Gland Tumor
H0159	Activated T-Cells, 8 hrs., ligation 2
H0163	Human Synovium
H0165	Human Prostate Cancer, Stage B2
H0166	Human Prostate Cancer, Stage B2 fraction
H0169	Human Prostate Cancer, Stage C fraction
H0170	12 Week Old Early Stage Human
H0171	12 Week Old Early Stage Human, II
H0173	Human Cardiomyopathy, RNA remake
H0176	CAMA1Ee Cell Line
H0177	CAMA1Ee Cell Line
H0178	Human Fetal Brain
H0179	Human Neutrophil
H0181	Human Primary Breast Cancer
H0182	Human Primary Breast Cancer
H0183	Human Colon Cancer
H0184	Human Colon Cancer, metasticized to live
H0186	Activated T-Cell
H0187	Resting T-Cell
H0188	Human Normal Breast
H0194	Human Cerebellum, subtracted
H0196	Human Cardiomyopathy, subtracted
H0197	Human Fetal Liver, subtracted
·	

770000	
H0200	Human Greater Omentum, fract II remake,
H0201	Human Hippocampus, subtracted
H0204	Human Colon Cancer, subtracted
H0205	Human Colon Cancer, differential
H0207	LNCAP, differential expression
H0208	Early Stage Human Lung, subtracted
H0212	Human Prostate, subtracted
H0213	Human Pituitary, subtracted
H0214	Raji cells, cyclohexamide treated, subtracted
H0216	Supt cells, cyclohexamide treated, subtracted
H0222	Activated T-Cells, 8 hrs, subtracted
H0225	Activated T-Cells, 12hrs, differentially expressed
H0231	Human Colon, subtraction
H0232	Human Colon, differential expression
H0234	human colon cancer, metastatic to liver, differentially expressed
H0235	Human colon cancer, metaticized to liver, subtraction
H0238	Human Myometrium Leiomyoma
H0239	Human Kidney Tumor
H0242	Human Fetal Heart, Differential (Fetal-Specific)
H0244	Human 8 Week Whole Embryo, subtracted
H0247	Human Membrane Bound Polysomes- Enzyme Subtraction
H0250	Human Activated Monocytes
H0251	Human Chondrosarcoma
H0252	Human Osteosarcoma
H0253	Human adult testis, large inserts
H0254	Breast Lymph node cDNA library
H0255	breast lymph node CDNA library
H0257	HL-60, PMA 4H
H0261	H. cerebellum, Enzyme subtracted
H0263	human colon cancer
H0264	human tonsils
H0265	Activated T-Cell (12hs)/Thiouridine labelledEco
H0266	Human Microvascular Endothelial Cells, fract. A
H0267	Human Microvascular Endothelial Cells, fract. B
H0268	Human Umbilical Vein Endothelial Cells, fract. A
H0269	Human Umbilical Vein Endothelial Cells, fract. B
H0271	Human Neutrophil, Activated
H0272	HUMAN TONSILS, FRACTION 2
H0274	Human Adult Spleen, fractionII
H0284	Human OB MG63 control fraction I
H0286	Human OB MG63 treated (10 nM E2) fraction I
H0288	Human OB HOS control fraction I
H0290	Human OB HOS treated (1 nM E2) fraction I
H0292	Human OB HOS treated (10 nM E2) fraction I
H0294	Amniotic Cells - TNF induced
H0295	Amniotic Cells - Primary Culture
H0298	HCBB's differential consolidation
H0305	CD34 positive cells (Cord Blood)
H0306	CD34 depleted Buffy Coat (Cord Blood)

H0309	Human Chronic Synovitis
H0313	human pleural cancer
H0316	HUMAN STOMACH
H0318	HUMAN B CELL LYMPHOMA
H0320	Human frontal cortex
H0321	HUMAN SCHWANOMA
H0327	human corpus colosum
H0328	human ovarian cancer
H0329	Dermatofibrosarcoma Protuberance
H0331	Hepatocellular Tumor
H0333	Hemangiopericytoma
H0334	Kidney cancer
H0340	Corpus Callosum
H0341	Bone Marrow Cell Line (RS4,11)
H0343	stomach cancer (human)
H0345	SKIN
H0351	Glioblastoma
H0352	wilm's tumor
H0354	Human Leukocytes
H0355	Human Liver
H0356	Human Kidney
H0359	KMH2 cell line
H0361	Human rejected kidney
H0369	H. Atrophic Endometrium
H0370	H. Lymph node breast Cancer
H0372	Human Testes
H0373	Human Heart
H0374	Human Brain
H0375	Human Lung
H0376	Human Spleen
H0379	Human Tongue, frac 1
H0380	Human Tongue, frac 2
H0381	Bone Cancer
H0383	Human Prostate BPH, re-excision
H0384	Brain, Kozak
H0386	Leukocyte and Lung, 4 screens
H0390	Human Amygdala Depression, re-excision
H0391	H. Meniingima, M6
H0392	H. Meningima, M1
H0393	Fetal Liver, subtraction II
H0395	A1-CELL LINE
H0399	Human Kidney Cortex, re-rescue
H0402	CD34 depleted Buffy Coat (Cord Blood), re-excision
H0403	H. Umbilical Vein Endothelial Cells, IL4 induced
H0408	Human kidney Cortex, subtracted
H0411	H Female Bladder, Adult
H0411	Human umbilical vein endothelial cells, IL-4 induced
H0413	Human Umbilical Vein Endothelial Cells, uninduced
H0415	H. Ovarian Tumor, II, OV5232
110713	11. Ovarian Tumor, 11, Ov 3232

Human Neutrophils, Activated, re-excision
Bone Cancer, re-excision
Human Bone Marrow, re-excision
T-Cell PHA 16 hrs
T-Cell PHA 24 hrs
Human Pituitary, subt IX
Human Adipose
Human Ovary
H. Kidney Medulla, re-excision
Human Umbilical Vein Endothelial cells, frac B, re-excision
Ovarian Tumor 10-3-95
Resting T-Cell Library,II
H Umbilical Vein Endothelial Cells, frac A, re-excision
H. Whole Brain #2, re-excision
H. Kidney Cortex, subtracted
Spleen metastic melanoma
Spleen, Chronic lymphocytic leukemia
H. Kidney Pyramid, subtracted
H. Striatum Depression, subt
Human Eosinophils
CD34+ cell, I, frac II
CD34+cells, II, FRACTION 2
H. Kidney Medulla, subtracted
Human Tonsil, Lib 3
Salivary Gland, Lib 2
Salivary Gland, Lib 3
Breast Cancer cell line, MDA 36
Breast Cancer Cell line, angiogenic
Hodgkin's Lymphoma I
Hodgkin's Lymphoma II
Human Tonsils, lib I
Human Tonsils, Lib 2
Crohn's Disease
HL-60, RA 4h, Subtracted
Keratinocyte
HEL cell line
Ulcerative Colitis
Liver, Hepatoma
Human Liver, normal
Keratinocyte, lib 3
Nasal polyps
pBMC stimulated w/ poly I/C
NTERA2, control
NTERA2 + retinoic acid, 14 days
Primary Dendritic Cells, lib 1
Primary Dendritic cells, frac 2
PCR, pBMC I/C treated
Poly[I]/Poly[C] Normal Lung Fibroblasts
Myoloid Progenitor Cell Line

H0530	Human Dermal Endothelial Cells,untreated
H0535	Human ovary tumor cell OV350721
H0538	Merkel Cells
H0539	Pancreas Islet Cell Tumor
H0540	Skin, burned
H0542	T Cell helper I
H0543	T cell helper II
H0544	Human endometrial stromal cells
H0545	Human endometrial stromal cells-treated with progesterone
H0546	Human endometrial stromal cells-treated with estradiol
H0547	NTERA2 teratocarcinoma cell line+retinoic acid (14 days)
H0549	H. Epididiymus, caput & corpus
H0550	H. Epididiymus, cauda
H0551	Human Thymus Stromal Cells
H0553	Human Placenta
H0555	Rejected Kidney, lib 4
H0556	Activated T-cell(12h)/Thiouridine-re-excision
H0559	HL-60, PMA 4H, re-excision
H0560	KMH2
H0561	L428
H0562	Human Fetal Brain, normalized c5-11-26
H0563	Human Fetal Brain, normalized 50021F
H0565	HUman Fetal Brain, normalized 100024F
H0569	Human Fetal Brain, normalized CO
H0570	Human Fetal Brain, normalized C500H
H0572	Human Fetal Brain, normalized AC5002
H0574	Hepatocellular Tumor, re-excision
H0575	Human Adult Pulmonary,re-excision
H0576	Resting T-Cell, re-excision
H0579	Pericardium
H0580	Dendritic cells, pooled
H0581	Human Bone Marrow, treated
H0583	B Cell lymphoma
H0584	Activated T-cells, 24 hrs,re-excision
H0585	Activated T-Cells, 12 hrs, re-excision
H0586	Healing groin wound, 6.5 hours post incision
H0587	Healing groin wound, 7.5 hours post incision
H0590	Human adult small intestine,re-excision
H0591	Human T-cell lymphoma,re-excision
H0592	Healing groin wound - zero hr post-incision (control)
H0593	Olfactory epithelium,nasalcavity
H0594	Human Lung Cancer,re-excision
H0595	Stomach cancer (human),re-excision
H0596_	Human Colon Cancer,re-excision
H0597	Human Colon, re-excision
H0598	Human Stomach,re-excision
H0599	Human Adult Heart,re-excision
H0600	Healing Abdomen wound,70&90 min post incision
H0601	Healing Abdomen Wound, 15 days post incision

	T
H0602	Healing Abdomen Wound,21&29 days post incision
H0606	Human Primary Breast Cancer,re-excision
H0609	H. Leukocytes, normalized cot > 500A
H0613	H.Leukocytes, normalized cot 5B
H0615	Human Ovarian Cancer Reexcision
H0616	Human Testes, Reexcision
H0617	Human Primary Breast Cancer Reexcision
H0618	Human Adult Testes, Large Inserts, Reexcision
H0619	Fetal Heart
H0620	Human Fetal Kidney, Reexcision
H0622	Human Pancreas Tumor, Reexcision
H0623	Human Umbilical Vein, Reexcision
H0624	12 Week Early Stage Human II, Reexcision
H0625	Ku 812F Basophils Line
H0626	Saos2 Cells, Untreated
H0627	Saos2 Cells, Vitamin D3 Treated
H0628	Human Pre-Differentiated Adipocytes
H0631	Saos2, Dexamethosome Treated
H0632	Hepatocellular Tumor,re-excision
H0633	Lung Carcinoma A549 TNFalpha activated
H0634	Human Testes Tumor, re-excision
H0635	Human Activated T-Cells, re-excision
H0638	CD40 activated monocyte dendridic cells
H0641	LPS activated derived dendritic cells
H0642	Hep G2 Cells, lambda library
H0643	Hep G2 Cells, PCR library
H0644	Human Placenta (re-excision)
H0645	Fetal Heart, re-excision
H0646	Lung, Cancer (4005313 A3): Invasive Poorly Differentiated Lung
	Adenocarcinoma,
H0647	Lung, Cancer (4005163 B7): Invasive, Poorly Diff. Adenocarcinoma, Metastatic
H0648	Ovary, Cancer: (4004562 B6) Papillary Serous Cystic Neoplasm, Low Malignant
	Pot
H0649	Lung, Normal: (4005313 B1)
H0650	B-Cells
H0651	Ovary, Normal: (9805C040R)
H0652	Lung, Normal: (4005313 B1)
H0653	Stromal Cells
H0654	Lung, Cancer: (4005313 A3) Invasive Poorly-differentiated Metastatic lung
	adenoc
H0656	B-cells (unstimulated)
H0657	B-cells (stimulated)
H0658	Ovary, Cancer (9809C332): Poorly differentiated adenocarcinoma
H0659	Ovary, Cancer (15395A1F): Grade II Papillary Carcinoma
H0660	Ovary, Cancer: (15799A1F) Poorly differentiated carcinoma
H0661	Breast, Cancer: (4004943 A5)
H0662	Breast, Normal: (4005522B2)
H0663	Breast, Cancer: (4005522 A2)
H0664	Breast, Cancer: (9806C012R)

LINGES	Stromal cells 3.88
H0665	
H0666	Ovary, Cancer: (4004332 A2) Stromal cells(HBM3.18)
H0667	stromal cells(HBM3.18)
H0668	
H0669	Breast, Cancer: (4005385 A2)
H0670	Ovary, Cancer(4004650 A3): Well-Differentiated Micropapillary Serous
110671	Carcinoma (0802C02OE)
H0671	Breast, Cancer: (9802C02OE)
H0672	Ovary, Cancer: (4004576 A8)
H0673	Human Prostate Cancer, Stage B2, re-excision
H0674	Human Prostate Cancer, Stage C, re-excission
H0675	Colon, Cancer: (9808C064R)
H0676	Colon, Cancer: (9808C064R)-total RNA
H0677	TNFR degenerate oligo
H0682	Ovarian cancer, Serous Papillary Adenocarcinoma
H0683	Ovarian Serous Papillary Adenocarcinoma
H0684	Serous Papillary Adenocarcinoma
H0685	Adenocarcinoma of Ovary, Human Cell Line, # OVCAR-3
H0686	Adenocarcinoma of Ovary, Human Cell Line
H0687	Human normal ovary(#9610G215)
H0688	Human Ovarian Cancer(#9807G017)
H0689	Ovarian Cancer
H0690	Ovarian Cancer, # 9702G001
H0691	Normal Ovary, #9710G208
H0693	Normal Prostate #ODQ3958EN
H0694	Prostate gland adenocarcinoma
H0695	mononucleocytes from patient
H0696	Prostate Adenocarcinoma
H0702	NK15(IL2 treated for 48 hours)
L0002	Atrium cDNA library Human heart
L0005	Clontech human aorta polyA+ mRNA (#6572)
L0021	Human adult (K.Okubo)
L0022	Human adult lung 3' directed MboI cDNA
L0040	Human colon mucosa
L0041	Human epidermal keratinocyte
L0053	Human pancreatic tumor
L0054	Human PGasparini
L0055	Human promyelocyte
L0065	Liver HepG2 cell line.
L0096	Subtracted human retina
L0105	Human aorta polyA+ (TFujiwara)
L0140	Human pancreatic cancer (CWallrapp)
L0142	Human placenta cDNA (TFujiwara)
L0143	Human placenta polyA+ (TFujiwara)
L0157	Human fetal brain (TFujiwara)
L0163	Human heart cDNA (YNakamura)
L0167	Human thymus (V.L.Boyartchuk)
L0193	Human osteosarcoma EGracia
L0194	Human pancreatic cancer cell line Patu 8988t

366

PCT/US00/26524

L0351	Infant brain, Bento Soares
L0351	Normalized infant brain, Bento Soares
L0352	Stratagene ovary (#937217)
L0362	
	Stratagene ovarian cancer (#937219)  NCI CGAP GC2
L0363	<del> </del>
L0364	NCI_CGAP_GC5
L0365	NCI_CGAP_Phe1
L0366	Stratagene schizo brain S11
L0367	NCI_CGAP_Sch1
L0368	NCI_CGAP_SS1
L0369	NCI_CGAP_AA1
L0370	Johnston frontal cortex
L0371	NCI_CGAP_Br3
L0372	NCI_CGAP_Co12
L0373	NCI_CGAP_Col1
L0374	NCI_CGAP_Co2
L0375	NCI CGAP Land
L0376	NCI_CGAP_Larl
L0378	NCI_CGAP_UDI4
L0381	NCI_CGAP_HN4
L0382	NCI CGAP Pr25
L0383	NCI_CGAP_Pr24
L0384	NCI_CGAP_Pr23
L0386	NCI_CGAP_HN3
L0387 L0388	NCI_CGAP_GCB0 NCI_CGAP_HN6
L0389	NCI CGAP HN5
L0389 L0394	H, Human adult Brain Cortex tissue
L0394 L0415	b4HB3MA Cot8-HAP-Ft
L0415	Infant brain, LLNL array of Dr. M. Soares 1NIB
L0433	normalized infant brain cDNA
L0439	Soares infant brain 1NIB
L0444	HB3MK
L0455	Human retina cDNA randomly primed sublibrary
L0456	Human retina cDNA Tsp509I-cleaved sublibrary
L0462	WATM1
L0471	Human fetal heart, Lambda ZAP Express
L0475	KG1-a Lambda Zap Express cDNA library
L0483	Human pancreatic islet
L0485	STRATAGENE Human skeletal muscle cDNA library, cat. #936215.
L0493	NCI CGAP Ov26
L0499	NCI CGAP HSC2
L0500	NCI CGAP Brn20
L0502	NCI CGAP Br15
L0503	NCI CGAP Br17
L0504	NCI CGAP Br13
L0505	NCI CGAP Br12
L0506	NCI CGAP Br16
L0507	NCI CGAP Br14
	1 * * * * * * * * * * * * * * * * * * *

	Large on the second
L0508	NCI_CGAP_Lu25
L0509	NCI_CGAP_Lu26
L0510	NCI_CGAP_Ov33
L0511	NCI_CGAP_Ov34
L0512	NCI_CGAP_Ov36
L0515	NCI_CGAP_Ov32
L0517	NCI_CGAP_Pr1
L0518	NCI_CGAP_Pr2
L0519	NCI_CGAP_Pr3
L0520	NCI_CGAP_Alv1
L0521	NCI_CGAP_Ew1
L0522	NCI_CGAP_Kid1
L0523	NCI_CGAP_Lip2
L0524	NCI_CGAP_Li1
L0526	NCI_CGAP_Pr12
L0527	NCI_CGAP_Ov2
L0528	NCI_CGAP_Pr5
L0529	NCI_CGAP_Pr6
L0530	NCI_CGAP_Pr8
L0532	NCI_CGAP_Thy1
L0534	Chromosome 7 Fetal Brain cDNA Library
L0539	Chromosome 7 Placental cDNA Library
L0540	NCI_CGAP_Pr10
L0541	NCI_CGAP_Pr7
L0542	NCI_CGAP_Prl1
L0543	NCI_CGAP_Pr9
L0544	NCI_CGAP_Pr4
L0545	NCI_CGAP_Pr4.1
L0553	NCI_CGAP_Co22
L0558	NCI_CGAP_Ov40
L0560	NCI_CGAP_HN12
L0562	Chromosome 7 HeLa cDNA Library
L0563	Human Bone Marrow Stromal Fibroblast
L0564	Jia bone marrow stroma
L0565	Normal Human Trabecular Bone Cells
L0581	Stratagene liver (#937224)
L0583	Stratagene cDNA library Human fibroblast, cat#937212
L0586	HTCDL1
L0587	Stratagene colon HT29 (#937221)
L0588	Stratagene endothelial cell 937223
L0589	Stratagene fetal retina 937202
L0590	Stratagene fibroblast (#937212)
L0591	Stratagene HeLa cell s3 937216
L0592	Stratagene hNT neuron (#937233)
L0593	Stratagene neuroepithelium (#937231)
L0594	Stratagene neuroepithelium NT2RAMI 937234
L0595	Stratagene NT2 neuronal precursor 937230
L0596	Stratagene colon (#937204)
L0597	Stratagene corneal stroma (#937222)

L0598	Morton Fetal Cochlea
L0599	Stratagene lung (#937210)
L0600	Weizmann Olfactory Epithelium
L0601	Stratagene pancreas (#937208)
L0602	Pancreatic Islet
L0603	Stratagene placenta (#937225)
L0604	Stratagene muscle 937209
L0605	Stratagene fetal spleen (#937205)
L0606	NCI_CGAP_Lym5
L0608	Stratagene lung carcinoma 937218
L0611	Schiller meningioma
L0617	Chromosome 22 exon
L0622	HM1
L0623	HM3
L0626	NCI_CGAP_GC1
L0627	NCI_CGAP_Co1
L0628	NCI_CGAP_Ov1
L0629	NCI CGAP Mel3
L0630	NCI_CGAP_CNS1
L0631	NCI CGAP Br7
L0634	NCI CGAP_Ov8
L0636	NCI CGAP Pit1
L0637	NCI CGAP Brn53
L0638	NCI CGAP Bm35
L0639	NCI CGAP Brn52
L0640	NCI CGAP_Br18
L0641	NCI CGAP Co17
L0642	NCI CGAP_Co18
L0643	NCI CGAP_Co19
L0644	NCI CGAP_Co20
L0645	NCI_CGAP_Co21
L0646	NCI CGAP_Co14
L0647	NCI CGAP Sar4
L0648	NCI CGAP_Eso2
L0649	NCI_CGAP_GU1
L0650	NCI_CGAP_Kid13
L0651	NCI_CGAP_Kid8
L0652	NCI_CGAP_Lu27
L0653	NCI_CGAP_Lu28
L0654	NCI CGAP_Lu31
L0655	NCI_CGAP_Lym12
L0656	NCI_CGAP_Ov38
L0657	NCI_CGAP_Ov23
L0658	NCI_CGAP_Ov35
L0659	NCI CGAP_Pan1
L0661	NCI_CGAP_Mel15
L0662	NCI CGAP_Gas4
L0663	NCI CGAP Ut2
L0664	NCI CGAP Ut3
L0656 L0657 L0658 L0659 L0661 L0662 L0663	NCI_CGAP_Ov38  NCI_CGAP_Ov23  NCI_CGAP_Ov35  NCI_CGAP_Pan1  NCI_CGAP_Mel15  NCI_CGAP_Gas4  NCI_CGAP_Ut2

L0665	NCI CGAP Ut4
L0666	NCI CGAP Utl
L0667	NCI CGAP CML1
L0683	Stanley Frontal NS pool 2
L0686	Stanley Frontal SN pool 2
L0689	Stanley Hippocampus SN pool 1
L0698	Testis 2
L0098	Gessler Wilms tumor
L0717	PN001-Normal Human Prostate
L0720	Soares pregnant uterus NbHPU
L0731 L0738	Human colorectal cancer
L0740	Soares melanocyte 2NbHM
L0741	Soares adult brain N2b4HB55Y
L0741	Soares adult brain N2b5HB55Y
L0742 L0743	
	Soares breast 2NbHBst
L0744	Soares breast 3NbHBst Soares retina N2b4HR
L0745	
L0746	Soares retina N2b5HR
L0747	Soares fetal heart NbHH19W
L0748	Soares fetal liver spleen 1NFLS
L0749	Soares_fetal_liver_spleen_1NFLS_S1
L0750 L0751	Soares_fetal_lung_NbHL19W  Soares_ovary_tumor_NbHOT
L0752	Soares_parathyroid_tumor_NbHPA
L0753	Soares_pineal_gland_N3HPG Soares placenta Nb2HP
L0754	Soares placenta 8to9weeks 2NbHP8to9W
L0755 L0756	Soares multiple_sclerosis_2NbHMSP
L0757	Soares senescent fibroblasts NbHSF
L0757	Soares testis NHT
L0759	Soares total fetus Nb2HF8 9w
L0759 L0761	NCI CGAP CLL1
L0761 L0762	NCI CGAP Br1.1
L0763	NCI CGAP Br2
L0764	NCI CGAP Co3
L0765	NCI CGAP_Co4
L0766	NCI CGAP GCB1
L0767	NCI CGAP GC3
L0767	NCI CGAP GC4
L0769	NCI CGAP Brn25
L0770	NCI CGAP Brn23
L0771	NCI CGAP Co8
L0772	NCI CGAP Col0
	NCI CGAP Co9
L0773	NCI CGAP Kid3
L0774	
L0775	NCL CGAP Lys
L0776	NCI_CGAP_Lu5
L0777	Soares_NhHMPu_S1
L0779	Soares_NFL_T_GBC_S1

L0780	Soares_NSF_F8_9W_OT_PA_P_S1
L0782	NCI_CGAP_Pr21
L0783	NCI_CGAP_Pr22
L0784	NCI CGAP_Lei2
L0785	Barstead spleen HPLRB2
L0786	Soares NbHFB
L0787	NCI CGAP_Sub1
L0788	NCI CGAP_Sub2
L0789	NCI CGAP_Sub3
L0790	NCI CGAP_Sub4
L0791	NCI_CGAP_Sub5
L0792	NCI CGAP_Sub6
L0793	NCI CGAP_Sub7
L0794	NCI CGAP_GC6
L0796	NCI CGAP_Brn50
L0800	NCI_CGAP_Co16
L0803	NCI_CGAP_Kid11
L0804	NCI_CGAP_Kid12
L0805	NCI_CGAP_Lu24
L0806	NCI_CGAP_Lu19
L0807	NCI CGAP Ov18
L0808	Barstead prostate BPH HPLRB4 1
L0809	NCI_CGAP_Pr28
N0007	Human Hippocampus
N0009	Human Hippocampus, prescreened
S0001	Brain frontal cortex
S0002	Monocyte activated
S0003	Human Osteoclastoma
S0004	Prostate
S0006	Neuroblastoma
S0007	Early Stage Human Brain
S0010	Human Amygdala
S0011	STROMAL -OSTEOCLASTOMA
S0013	Prostate
S0014	Kidney Cortex
S0015	Kidney medulla
S0022	Human Osteoclastoma Stromal Cells - unamplified
S0026	Stromal cell TF274
S0027	Smooth muscle, serum treated
S0028	Smooth muscle,control
S0029	brain stem
S0031	Spinal cord
S0032	Smooth muscle-ILb induced
S0036	Human Substantia Nigra
S0037	Smooth muscle, IL1b induced
S0038	Human Whole Brain #2 - Oligo dT > 1.5Kb
S0040	Adipocytes
S0044	Prostate BPH
S0045	Endothelial cells-control

S0046	Endothelial-induced
S0048	Human Hypothalamus, Alzheimer's
S0049	Human Brain, Striatum
S0050	Human Frontal Cortex, Schizophrenia
S0051	Human Hypothalmus, Schizophrenia
S0052	neutrophils control
S0053	Neutrophils IL-1 and LPS induced
S0106	STRIATUM DEPRESSION
S0112	Hypothalamus
S0114	Anergic T-cell
S0116	Bone marrow
S0122	Osteoclastoma-normalized A
S0126	Osteoblasts
S0132	Epithelial-TNFa and INF induced
S0134	Apoptotic T-cell
S0136	PERM TF274
S0140	eosinophil-IL5 induced
S0142	Macrophage-oxLDL
S0144	Macrophage (GM-CSF treated)
S0146	prostate-edited
S0148	Normal Prostate
S0150	LNCAP prostate cell line
S0152	PC3 Prostate cell line
S0174	Prostate-BPH subtracted II
S0176	Prostate, normal, subtraction I
S0182	Human B Cell 8866
S0188	Prostate, BPH, Lib 2
S0190	Prostate BPH,Lib 2, subtracted
S0192	Synovial Fibroblasts (control)
S0194	Synovial hypoxia
S0196	Synovial IL-1/TNF stimulated
S0206	Smooth Muscle- HASTE normalized
S0208	Messangial cell, frac 1
S0210	Messangial cell, frac 2
S0212	Bone Marrow Stromal Cell, untreated
S0214	Human Osteoclastoma, re-excision
S0216	Neutrophils IL-1 and LPS induced
S0218	Apoptotic T-cell, re-excision
S0222	H. Frontal cortex, epileptic, re-excision
S0242	Synovial Fibroblasts (II1/TNF), subt
S0250	Human Osteoblasts II
S0260	Spinal Cord, re-excision
S0276	Synovial hypoxia-RSF subtracted
S0278	H Macrophage (GM-CSF treated), re-excision
S0280	Human Adipose Tissue, re-excision
S0282	Brain Frontal Cortex, re-excision
S0294	Larynx tumor
S0300	Frontal lobe,dementia,re-excision
S0306	Larynx normal #10 261-273
· · · · · · · · · · · · · · · · · · ·	

S0312	Human osteoarthritic, fraction II
S0314	Human osteoarthritis, fraction I
S0318	Human Normal Cartilage Fraction II
S0320	Human Larynx
S0322	Siebben Polyposis
S0328	Palate carcinoma
S0330	Palate normal
S0332	Pharynx carcinoma
S0334	Human Normal Cartilage Fraction III
S0342	Adipocytes,re-excision
S0344	Macrophage-oxLDL, re-excision
S0346	Human Amygdala,re-excision
S0348	Cheek Carcinoma
S0350	Pharynx Carcinoma
S0354	Colon Normal II
S0356	Colon Carcinoma
S0358	Colon Normal III
S0360	Colon Tumor II
S0362	Human Gastrocnemius
S0364	Human Quadriceps
S0366	Human Soleus
S0370	Larynx carcinoma II
S0372	Larynx carcinoma III
S0374	Normal colon
S0376	Colon Tumor
S0378	Pancreas normal PCA4 No
S0380	Pancreas Tumor PCA4 Tu
S0382	Larynx carcinoma IV
S0384	Tongue carcinoma
S0386	Human Whole Brain, re-excision
S0388	Human Hypothalamus, schizophrenia, re-excision
S0390	Smooth muscle, control, re-excision
S0392	Salivary Gland
S0394	Stomach,normal
S0402	Adrenal Gland,normal
S0404	Rectum normal
S0406	Rectum tumour
S0408	Colon, normal
S0410	Colon, tumour
S0414	Hippocampus, Alzheimer Subtracted
S0418	CHME Cell Line,treated 5 hrs
S0420	CHME Cell Line,untreated
S0422	Mo7e Cell Line GM-CSF treated (1ng/ml)
S0424	TF-1 Cell Line GM-CSF Treated
S0426	Monocyte activated, re-excision
S0428	Neutrophils control, re-excision
S0430	Aryepiglottis Normal
S0432	Sinus piniformis Tumour
S0434	Stomach Normal

S0440		
S0440	S0436	Stomach Tumour
S0442   Colon Normal	S0438	
S0444	S0440	Liver Tumour Met 5 Tu
S0446	S0442	Colon Normal
S0448	S0444	Colon Tumor
S0450	S0446	Tongue Tumour
S0452   Thymus	S0448	Larynx Normal
S0454   Placenta   S0456   Tongue Normal   S0458   Thyroid Normal (SDCA2 No)   S0460   Thyroid Tumour   S0462   Thyroid Thyroiditis   S0464   Larynx Normal   S0464   Larynx Normal   S0464   Larynx Normal   S0465   Ea.hy.926 cell line   S0472   Lung Mesothelium   S0474   Human blood platelets   S3012   Smooth Muscle Serum Treated, Norm   S3014   Smooth muscle, serum induced,re-exc   S6014   H. hypothalamus, frac A   S6016   H. Frontal Cortex, Epileptic   S6022   H. Adipose Tissue   S6024   Alzheimers, spongy change   S6026   Frontal Lobe, Dementia   S6028   Human Manic Depression Tissue   T0002   Activated T-cells   T0003   Human Fetal Lung   T0006   Human Pincal Gland   T0008   Colorectal Tumor   T0010   Human Infant Brain   T0023   Human Pancreatic Carcinoma   T0039   HSA 172 Cells   T0041   Jurkat T-cell G1 phase   T0042   Jurkat T-cell G1 phase   T0043   Human Adult Retina   T0066   Human Thyroid   T0067   Human Thyroid   T0068   Normal Ovary, Premenopausal   T0069   Human Uterus, normal   T0071   Human Bone Marrow   Human Adult Retina   T0079   Human Kidney, normal Adult   T0082   Human Adult Retina   T0079   Human Kidney, normal Adult   T0082   Human Adult Retina   T0082   Human	S0450	Larynx Tumour
S0456	S0452	Thymus
S0458   Thyroid Normal (SDCA2 No)	S0454	Placenta
S0460   Thyroid Tumour	S0456	Tongue Normal
S0462 Thyroid Thyroiditis S0464 Larynx Normal S0468 Ea.hy.926 cell line S0472 Lung Mesothelium S0474 Human blood platelets S3012 Smooth Muscle Serum Treated, Norm S3014 Smooth muscle, serum induced,re-exc S6014 H. hypothalamus, frac A S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0043 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult	S0458	Thyroid Normal (SDCA2 No)
S0464 Larynx Normal S0468 Ea.hy.926 cell line S0472 Lung Mesothelium S0474 Human blood platelets S3012 Smooth Muscle Serum Treated, Norm S3014 Smooth muscle, serum induced,re-exc S6014 H. hypothalamus, frac A S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0043 Human Aortic Endothelium T0044 Aorta endothelial cells + TNF-a T0066 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult	S0460	Thyroid Tumour
S0468 Ea.hy.926 cell line S0472 Lung Mesothelium S0474 Human blood platelets S3012 Smooth Muscle Serum Treated, Norm S3014 Smooth muscle, serum induced,re-exc S6014 H. hypothalamus, frac A S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell Gl phase T0042 Jurkat T-cell Gl phase T0043 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult	S0462	Thyroid Thyroiditis
S0472 Lung Mesothelium S0474 Human blood platelets S3012 Smooth Muscle Serum Treated, Norm S3014 Smooth muscle, serum induced,re-exc S6014 H. hypothalamus, frac A S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0043 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult	S0464	Larynx Normal
S0474 Human blood platelets S3012 Smooth Muscle Serum Treated, Norm S3014 Smooth muscle, serum induced,re-exc S6014 H. hypothalamus, frac A S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0043 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Bone Marrow T0074 Human Bone Marrow T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0082 Human Kidney, normal Adult	S0468	Ea.hy.926 cell line
S3012 Smooth Muscle Serum Treated, Norm S3014 Smooth muscle, serum induced, re-exc S6014 H. hypothalamus, frac A S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell GI phase T0042 Jurkat T-Cell, S phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0082 Human Kidney, normal Adult	S0472	Lung Mesothelium
S3014 Smooth muscle, serum induced, re-exc S6014 H. hypothalamus, frac A S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0082 Human Kidney, normal Adult	S0474	Human blood platelets
S6014 H. hypothalamus, frac A S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0043 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult	S3012	Smooth Muscle Serum Treated, Norm
S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0082 Human Adult Retina	S3014	Smooth muscle, serum induced,re-exc
S6016 H. Frontal Cortex, Epileptic S6022 H. Adipose Tissue S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0082 Human Adult Retina	S6014	H. hypothalamus, frac A
S6024 Alzheimers, spongy change S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0070 Human Kidney, normal Adult T0070 Human Kidney, normal Adult	S6016	
S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-cell G1 phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0082 Human Adult Retina	S6022	H. Adipose Tissue
S6026 Frontal Lobe, Dementia S6028 Human Manic Depression Tissue T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-cell G1 phase T0043 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0079 Human Kidney, normal Adult T0079 Human Adult Retina	S6024	Alzheimers, spongy change
T0002 Activated T-cells T0003 Human Fetal Lung T0006 Human Pincal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	S6026	
T0003 Human Fetal Lung T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell Gl phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0070 Human Kidney, normal Adult T0070 Human Kidney, normal Adult T0070 Human Adult Retina	S6028	Human Manic Depression Tissue
T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0002	Activated T-cells
T0006 Human Pineal Gland T0008 Colorectal Tumor T0010 Human Infant Brain T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0003	Human Fetal Lung
T0010 Human Infant Brain  T0023 Human Pancreatic Carcinoma  T0039 HSA 172 Cells  T0040 HSC172 cells  T0041 Jurkat T-cell G1 phase  T0042 Jurkat T-Cell, S phase  T0048 Human Aortic Endothelium  T0049 Aorta endothelial cells + TNF-a  T0060 Human White Adipose  T0067 Human Thyroid  T0068 Normal Ovary, Premenopausal  T0069 Human Uterus, normal  T0071 Human Bone Marrow  T0074 Human Adult Retina  T0079 Human Kidney, normal Adult  T0082 Human Adult Retina	T0006	
T0023 Human Pancreatic Carcinoma T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0008	Colorectal Tumor
T0039 HSA 172 Cells T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0010	Human Infant Brain
T0040 HSC172 cells T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0023	Human Pancreatic Carcinoma
T0041 Jurkat T-cell G1 phase T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0039	HSA 172 Cells
T0042 Jurkat T-Cell, S phase T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0040	HSC172 cells
T0048 Human Aortic Endothelium T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0041	Jurkat T-cell G1 phase
T0049 Aorta endothelial cells + TNF-a T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0042	Jurkat T-Cell, S phase
T0060 Human White Adipose T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0048	Human Aortic Endothelium
T0067 Human Thyroid T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0049	Aorta endothelial cells + TNF-a
T0068 Normal Ovary, Premenopausal T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0060	Human White Adipose
T0069 Human Uterus, normal T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0067	Human Thyroid
T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0068	Normal Ovary, Premenopausal
T0071 Human Bone Marrow T0074 Human Adult Retina T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0069	Human Uterus, normal
T0079 Human Kidney, normal Adult T0082 Human Adult Retina	T0071	
T0082 Human Adult Retina	T0074	Human Adult Retina
T0082 Human Adult Retina	T0079	Human Kidney, normal Adult
T0102 Human colon carainoma (HCC) cell line		
10103   Human colon caremonia (ACC) cen line	T0103	Human colon carcinoma (HCC) cell line
T0104 HCC cell line metastisis to liver	T0104	<del>                                     </del>
T0109 Human (HCC) cell line liver (mouse) metastasis, remake	T0109	Human (HCC) cell line liver (mouse) metastasis, remake

T0110	Human colon carcinoma (HCC) cell line, remake
T0114	Human (Caco-2) cell line, adenocarcinoma, colon, remake
T0115	Human Colon Carcinoma (HCC) cell line

Table 6

MIM ID	OMIM Description
100650	Alcohol intolerance, acute (3)
	?Fetal alcohol syndrome (1)
100690	Myasthenic syndrome, slow-channel congenital, 601462 (3)
100710	Myasthenic syndrome, slow-channel congenital, 601462 (3)
100730	Myasthenia gravis, neonatal transient (2)
101000	Malignant mesothelioma, sporadic (3)
	Meningioma, NF2-related, sporadic (3) Schwannoma, sporadic (3)
	Neurofibromatosis, type 2 (3)
	Neurolemmomatosis (3)
102200	Somatotrophinoma (2)
102540	Cardiomyopathy, idiopathic dilated (3)
102578	Leukemia, acute promyelocytic, PML/RARA type (3)
102600	Urolithiasis, 2,8-dihydroxyadenine (3)
102770	Myoadenylate deaminase deficiency (3)
102772	[AMP deaminase deficiency, erythrocytic] (3)
103000	Hemolytic anemia due to adenylate kinase deficiency (3)
103050	Adenylosuccinase deficiency (1)
	Autism, succinylpurinemic (3)
103581	Albright hereditary osteodystrophy-2 (2) (?)
103600	Analbuminemia (3)
	[Dysalbuminemic hyperthyroxinemia] (3)
	[Dysalbuminemic hyperzincemia], 194470 (3)
103720	Alcoholism, susceptibility to (1)
103850	Aldolase A deficiency (3)
103950	Emphysema due to alpha-2-macroglobulin deficiency (1)
104150	[AFP deficiency, congenital] (1)
	[Hereditary persistence of alpha-fetoprotein] (3)
104311	Alzheimer disease-3 (3)
104500	Amelogenesis imperfecta-2, hypoplastic local type (2)
104614	Cystinuria, 220100 (3)
104770	?Amyloidosis, secondary, susceptibility to (1)
105580	Anal canal carcinoma (2) (?)
105600	Dyserythropoietic anemia, congenital, type III (2)
106100	Angioedema, hereditary (3)
106150	Hypertension, essential, susceptibility to (3)
	Preeclampsia, susceptibility to (3)
106165	Hypertension, essential, 145500 (3)
106180	Myocardial infarction, susceptibility to (3)
106210	Aniridia (3)
	Cataract, congenital, with late-onset corneal dystrophy (3)
	Foveal hypoplasia, isolated, 136520 (3)
	Peters anomaly (3)
106300	Ankylosing spondylitis (2)
107250	Anterior segment mesenchymal dysgenesis (2)
107271	CD59 deficiency (3)
107280	Alpha-1-antichymotrypsin deficiency (3)
	Cerebrovascular disease, occlusive (3)

107300	Antithrombin III deficiency (3)
107400	Emphysema (3)
107100	Emphysema-cirrhosis (3)
107470	Atypical mycobacterial infection, familial disseminated, 209950 (3)
107170	BCG infection, generalized familial (3)
	Tuberculosis, susceptibility to (3)
107670	Apolipoprotein A-II deficiency (3)
107680	Amyloidosis, 3 or more types (3)
	ApoA-I and apoC-III deficiency, combined (3)
	Corneal clouding, autosomal recessive (3)
	Hypertriglyceridemia, one form (3)
	Hypoalphalipoproteinemia (3)
107720	Hypertriglyceridemia (3)
107730	Abetalipoproteinemia (3)
	Apolipoprotein B-100, ligand-defective (3)
	Hyperbetalipoproteinemia (3)
	Hypobetalipoproteinemia (3)
107741	Hyperlipoproteinemia, type III (3)
107776	Colton blood group, 110450 (3)
107777	Diabetes insipidus, nephrogenic, autosomal recessive, 222000 (3)
107910	Gynecomastia, familial, due to increased aromatase activity (1)
	Virilization, maternal and fetal, from placental aromatase deficiency (3)
107970	Arrhythmogenic right ventricular dysplasia-1 (2)
108120	Distal arthrogryposis-1 (2)
108725	Atherosclerosis, susceptibility to (2)
108730	Brody myopathy, 601003 (3)
108800	Atrial septal defect, secundum type (2)
108962	Hypertension, salt-resistant (1) (?)
108985	Atrophia areata (2)
109150	Machado-Joseph disease (3)
109270	Hemolytic anemia due to band 3 defect (3)
	Renal tubular acidosis, distal, 179800 (3)
	Spherocytosis, hereditary (3)
	[Acanthocytosis, one form] (1)
100100	[Elliptocytosis, Malaysian-Melanesian type] (3)
109400	Basal cell nevus syndrome (2)
109543	Leukemia, chronic lymphocytic, B-cell (2)
109560	Leukemia/lymphoma, B-cell, 3 (2)
109690	Asthma, nocturnal, susceptibility to (3)
100700	Obesity, susceptibility to (3)
109700	Hemodialysis-related amyloidosis (1)
110100	Blepharophimosis, epicanthus inversus, and ptosis, type 1 (2)
110700	Vivax malaria, susceptibility to (1)
112250	Bone dysplasia with medullary fibrosarcoma (2)
112261	Fibrodysplasia ossificans progressiva (1) (?)
112262	Fibrodysplasia ossificans progressiva, 135100 (1) (?)
112410	Hypertension with brachydactyly (2)
113100	Brachydactyly, type C (2)
113300	Brachydactyly type E (2) (?)
113520	Hyperleucinemia-isoleucinemia or hypervalinemia (1) (?)

113705	Breast cancer-1 (3)
113703	Ovarian cancer (3)
113721	Breast cancer (1)
113900	Heart block, progressive familial, type I (2)
114130	Osteoporosis (3)
114208	Hypokalemic periodic paralysis, 170400 (3)
114206	Malignant hyperthermia susceptibility 5, 601887 (3)
114240	Muscular dystrophy, limb-girdle, type 2A, 253600 (3)
114290	Campomelic dysplasia with autosomal sex reversal (3)
	Leukemia, acute myeloid (2)
114350	
114400	Lynch cancer family syndrome II (2) (?)
114550	Hepatocellular carcinoma (1)
114835	Monocyte carboxyesterase deficiency (1) (?)
115500	Acatalasemia (3)
115650	Cataract, anterior polar-1 (2) (?)
115660	Cataract, cerulean, type 1 (2)
116600	Cataract, posterior polar (2)
116800	Cataract, Marner type (2)
116806	Colorectal cancer (3)
116860	Cavernous angiomatous malformations (2)
117700	Hemosiderosis, systemic, due to aceruloplasminemia (3)
•	[Hypoceruloplasminemia, hereditary] (1)
118210	Charcot-Marie-Tooth neuropathy-2A (2)
118425	Myotonia congenita, dominant, 160800 (3)
	Myotonia congenita, recessive, 255700 (3)
	Myotonia levior, recessive (3)
118470	[CETP deficiency] (3)
118485	Polycystic ovary syndrome with hyperandrogenemia (2)
118504	Epilepsy, benign neonatal, type 1, 121200 (3)
	Epilepsy, nocturnal frontal lobe, 600513 (3)
118511	Schizophrenia, neurophysiologic defect in (2)
118800	Choreoathetosis, familial paroxysmal (2)
119300	van der Woude syndrome (2)
120070	Alport syndrome, autosomal recessive, 203780 (3)
120110	Metaphyseal chondrodysplasia, Schmid type (3)
120120	Epidermolysis bullosa dystrophica, dominant, 131750 (3)
	Epidermolysis bullosa dystrophica, recessive, 226600 (3)
	Epidermolysis bullosa, pretibial, 131850 (3)
120131	Alport syndrome, autosomal recessive, 203780 (3)
	Hematuria, familial benign (3)
120140	Achondrogenesis-hypochondrogenesis, type II (3)
	Kniest dysplasia (3)
	Osteoarthrosis, precocious (3)
	SED congenita (3)
	SMED Strudwick type (3)
	Stickler syndrome, type I (3)
	Wagner syndrome, type II (3)
120150	Ehlers-Danlos syndrome, type VIIA1, 130060 (3)
	Osteogenesis imperfecta, 4 clinical forms, 166200, 166210, 259420,
	166220 (3)

Osteoporosis, idiopathic, 166710 (3)
Ehlers-Danlos syndrome, type VIIA2, 130060 (3)
Marfan syndrome, atypical (3)
Osteogenesis imperfecta, 4 clinical forms, 166200, 166210, 259420,
166220 (3)
Osteoporosis, idiopathic, 166710 (3)
Aneurysm, familial, 100070 (3)
Ehlers-Danlos syndrome, type III (3)
Ehlers-Danlos syndrome, type IV, 130050 (3)
Fibromuscular dysplasia of arteries, 135580 (3)
Ehlers-Danlos syndrome, type I, 130000 (3)
Ehlers-Danlos syndrome, type I, 130000 (3)
Ehlers-Danlos syndrome, type II, 130010 (3)
Bethlem myopathy, 158810 (3)
Bethlem myopathy, 158810 (3)
Bethlem myopathy, 158810 (3)
Epiphyseal dysplasia, multiple, type 2, 600204 (3)
Marshall syndrome, 154780 (3)
Stickler syndrome, type III (3)
OSMED syndrome, 215150 (3)
Stickler syndrome, type II, 184840 (3)
Colorectal cancer, hereditary, nonpolyposis, type 1 (3) Ovarian cancer (3)
Muir-Torre syndrome, 158320 (3)
The second secon
Colorectal cancer, hereditary nonpolyposis, type 2 (3)
Muir-Torre family cancer syndrome, 158320 (3)
Turcot syndrome with glioblastoma, 276300 (3)
Colorectal cancer (3)
Clq deficiency, type A (3)
Clq deficiency, type B (3)
Clq deficiency, type C (3)
C1r/C1s deficiency, combined (1)
CR1 deficiency (1)
1.781 b cuccentibility (1)
?SLE susceptibility (1)
C3 deficiency (3)
C3 deficiency (3) C4 deficiency (3)
C3 deficiency (3) C4 deficiency (3) C4 deficiency (3)
C3 deficiency (3) C4 deficiency (3) C4 deficiency (3) C5 deficiency (1)
C3 deficiency (3) C4 deficiency (3) C4 deficiency (3) C5 deficiency (1) Measles, susceptibility to (1)
C3 deficiency (3) C4 deficiency (3) C4 deficiency (3) C5 deficiency (1) Measles, susceptibility to (1) C9 deficiency (3)
C3 deficiency (3) C4 deficiency (3) C4 deficiency (3) C5 deficiency (1) Measles, susceptibility to (1) C9 deficiency (3) C8 deficiency, type I (2)
C3 deficiency (3) C4 deficiency (3) C4 deficiency (3) C5 deficiency (1) Measles, susceptibility to (1) C9 deficiency (3) C8 deficiency, type I (2) C8 deficiency, type II (3)
C3 deficiency (3)  C4 deficiency (3)  C4 deficiency (3)  C5 deficiency (1)  Measles, susceptibility to (1)  C9 deficiency (3)  C8 deficiency, type I (2)  C8 deficiency, type II (3)  Deafness, autosomal dominant 3, 601544 (3)
C3 deficiency (3) C4 deficiency (3) C4 deficiency (3) C5 deficiency (1) Measles, susceptibility to (1) C9 deficiency (3) C8 deficiency, type I (2) C8 deficiency, type II (3)
C3 deficiency (3)  C4 deficiency (3)  C4 deficiency (3)  C5 deficiency (1)  Measles, susceptibility to (1)  C9 deficiency (3)  C8 deficiency, type I (2)  C8 deficiency, type II (3)  Deafness, autosomal dominant 3, 601544 (3)
C3 deficiency (3) C4 deficiency (3) C5 deficiency (1) Measles, susceptibility to (1) C9 deficiency (3) C8 deficiency, type I (2) C8 deficiency, type II (3) Deafness, autosomal dominant 3, 601544 (3) Deafness, autosomal recessive 1, 220290 (3)
C3 deficiency (3) C4 deficiency (3) C5 deficiency (1) Measles, susceptibility to (1) C9 deficiency (3) C8 deficiency, type I (2) C8 deficiency, type II (3) Deafness, autosomal dominant 3, 601544 (3) Deafness, autosomal recessive 1, 220290 (3) Heterotaxia, visceroatrial, autosomal recessive (3)
C3 deficiency (3) C4 deficiency (3) C5 deficiency (1) Measles, susceptibility to (1) C9 deficiency (3) C8 deficiency, type I (2) C8 deficiency, type II (3) Deafness, autosomal dominant 3, 601544 (3) Deafness, autosomal recessive 1, 220290 (3) Heterotaxia, visceroatrial, autosomal recessive (3) Contractural arachnodactyly, congenital (3)
C3 deficiency (3)  C4 deficiency (3)  C5 deficiency (1)  Measles, susceptibility to (1)  C9 deficiency (3)  C8 deficiency, type I (2)  C8 deficiency, type II (3)  Deafness, autosomal dominant 3, 601544 (3)  Deafness, autosomal recessive 1, 220290 (3)  Heterotaxia, visceroatrial, autosomal recessive (3)  Contractural arachnodactyly, congenital (3)  Coproporphyria (3)
C3 deficiency (3)  C4 deficiency (3)  C5 deficiency (1)  Measles, susceptibility to (1)  C9 deficiency (3)  C8 deficiency, type I (2)  C8 deficiency, type II (3)  Deafness, autosomal dominant 3, 601544 (3)  Deafness, autosomal recessive 1, 220290 (3)  Heterotaxia, visceroatrial, autosomal recessive (3)  Contractural arachnodactyly, congenital (3)  Coproporphyria (3)  Harderoporphyrinuria (3)

122000	Corneal dystrophy, posterior polymorphous (2)
122500	[Transcortin deficiency] (1) ACTH deficiency, 201400 (2)
122720	Coumarin resistance, 122700 (3)
122000	Nicotine addiction, protection from (3)
123000	Craniometaphyseal dysplasia (2)
123101	Craniosynostosis, type 2 (3)
123580	Cataract, congenital, autosomal dominant (3)
123620	Cataract, cerulean, type 2, 601547 (3)
123660	Cataract, Coppock-like (3)
123829	Melanoma (3)
123940	White sponge nevus, 193900 (3)
124030	Debrisoquine sensitivity (3)
	?Parkinsonism, susceptibility to (1)
124080	CMO II deficiency (3)
124200	Darier disease (keratosis follicularis) (2)
125270	Porphyria, acute hepatic (3)
	Lead poisoning, susceptibility to (3)
125490	Dentinogenesis imperfecta-1 (2)
125660	Cardiomyopathy (1) (?)
	Myopathy, desminopathic (1) (?)
125852	Insulin-dependent diabetes mellitus-2 (2)
126060	Anemia, megaloblastic, due to DHFR deficiency (1) (?)
126090	Hyperphenylalaninemia due to pterin-4a-carbinolamine dehydratase deficiency, 264070 (3)
126150	Diphtheria, susceptibility to (1)
126337	Myxoid liposarcoma (3)
126340	Xeroderma pigmentosum, group D, 278730 (3)
126391	DNA ligase I deficiency (3)
126451	?Schizophrenia, susceptibility to (2)
126452	Autonomic nervous system dysfunction (3)
120.02	[Novelty seeking personality] (1)
126600	Drusen, radial, autosomal dominant (2)
126650	Chloride diarrhea, congenital, Finnish type, 214700 (3)
12000	Colon cancer (1) (?)
128100	Dystonia-1, torsion (3)
129010	Neuropathy, congenital hypomyelinating, 1 (3)
129490	Ectodermal dysplasia-3, anhidrotic (2)
129500	Ectodermal dysplasia, hidrotic (2)
129900	EEC syndrome-1 (2) (?)
130160	Cutis laxa, 123700 (3)
	Supravalvar aortic stenosis, 185500 (3)
	Williams-Beuren syndrome, 194050 (3)
130410	Glutaricaciduria, type IIB (3)
130500	Elliptocytosis-1 (3)
130650	Beckwith-Wiedemann syndrome (2)
131100	Carcinoid tumor of lung (3)
	Multiple endocrine neoplasia I (3)
	Prolactinoma, hyperparathyroidism, carcinoid syndrome (2)
L	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -

131195	Hereditary hemorrhagic telangiectasia-1, 187300 (3)
131210	Atherosclerosis, susceptibility to (2)
131242	Shah-Waardenburg syndrome, 277580 (3)
131400	Eosinophilia, familial (2)
131440	Eosinophilic myeloproliferative disorder (2) (?)
132700	Cylindromatosis (2)
132800	Basal cell carcinoma (2) (?)
132600	Epithelioma, self-healing, squamous 1, Ferguson-Smith type (2)
133170	Erythremia (1) (?)
133171	[Erythrocytosis, familial], 133100 (3)
133200	Erythrokeratodermia variabilis (2)
133510	Trichothiodystrophy (3)
100010	Xeroderma pigmentosum, group B (3)
133530	Xeroderma pigmentosum, group G, 278780 (3)
133540	Cockayne syndrome-2, late onset (2)
133700	Chondrosarcoma, 215300 (3)
	Exostoses, multiple, type 1 (3)
133701	Exostoses, multiple, type 2 (3)
133780	Vitreoretinopathy, exudative, familial (2)
134370	Factor H deficiency (1)
	Hemolytic-uremic syndrome, 235400 (3)
	Membroproliferative glomerulonephritis (1)
134580	Factor XIIIB deficiency (3)
134638	Systemic lupus erythematosus, susceptibility, 152700 (3)
134790	Hyperferritinemia-cataract syndrome, 600886 (3)
134797	Ectopia lentis, ?isolated (3)
	Marfan syndrome, 154700 (3)
	Shprintzen-Goldberg syndrome, 182212 (3)
134820	Amyloidosis, hereditary renal, 105200 (3)
	Dysfibrinogenemia, alpha type, causing bleeding diathesis (3)
	Dysfibrinogenemia, alpha type, causing recurrent thrombosis (3)
134830	Dysfibrinogenemia, beta type (3)
134850	Dysfibrinogenemia, gamma type (3)
	Hypofibrinogenemia, gamma type (3)
134934	Achondroplasia, 100800 (3)
	Craniosynostosis, nonsyndromic (3)
	Crouzon syndrome with acanthosis nigricans (3)
	Hypochondroplasia, 146000 (3)
105000	Thanatophoric dysplasia, types I and II, 187600 (3)
135300	Fibromatosis, gingival (2)
135600	Ehlers-Danlos syndrome, type X (1) (?)
135700	Fibrosis of extraocular muscles, congenital, 1 (2)
135750	Tetramelic mirror-image polydactyly (2) (?)
135940	Ichthyosis vulgaris, 146700 (1) (?)
136132	[Fish-odor syndrome], 602079 (3)
136350	Pfeiffer syndrome, 101600 (3)
136435	Ovarian dysgenesis, hypergonadotropic, with normal karyotype, 233300 (3)
136440	Lymphoma/leukemia, B-cell, variant (1)
136530	Male infertility, familial (1) (?)
136550	Macular dystrophy, North Carolina type (2)

136836	Fucosyltransferase-6 deficiency (3)
136850	Fumarase deficiency (3)
137181	[Gamma-glutamyltransferase, familial high serum] (2)
137350	Amyloidosis, Finnish type, 105120 (3)
137600	Iridogoniodysgenesis syndrome (2)
138030	[?Hyperproglucagonemia] (1)
138033	Diabetes mellitus, type II (3)
138040	Cortisol resistance (3)
138079	Hyperinsulinism, familial, 602485 (3) MODY, type 2, 125851 (3)
138140	Glucose transport defect, blood-brain barrier (3)
138190	Diabetes mellitus, noninsulin-dependent (3)
138320	Hemolytic anemia due to glutathione peroxidase deficiency (1)
138430	Diabetes mellitus, type II (3)
138491	Hyperekplexia and spastic paraparesis (3)
150451	Startle disease, autosomal recessive (3)
	Startle disease/hyperekplexia, autosomal dominant, 149400 (3)
138570	Non-insulin dependent diabetes mellitus, susceptibility to (2)
138571	Glycogen synthase, liver, deficiency of, 240600 (1)
138700	[Apolipoprotein H deficiency] (3)
138720	Bernard-Soulier syndrome, type B (2)
138850	Hypogonadotropic hypogonadism (3)
138971	Kostmann neutropenia, 202700 (3)
138981	Pulmonary alveolar proteinosis, 265120 (3)
139130	Hypertension, essential, susceptibility to, 145500 (3)
	Basal cell carcinoma (3)
139150	Gigantism due to GHRF hypersecretion (1)
139190	Isolated growth hormone deficiency due to defect in GHRF (1) (?)
139191	Growth hormone deficient dwarfism (3)
	Isolated growth hormone deficiency, Illig type with absent GH and
139250	Kowarski type with bioinactive GH (3)
139320	McCune-Albright polyostotic fibrous dysplasia, 174800 (3)
	Pituitary ACTH secreting adenoma (3)
	Pseudohypoparathyroidism, type Ia, 103580 (3)
	Somatotrophinoma (3)
139330	Night blindness, congenital stationary (3)
139350	Epidermolytic hyperkeratosis, 113800 (3)
	Keratoderma, palmoplantar, nonepidermolytic (3)
139360	Pituitary ACTH-secreting adenoma (3)
140100	[Anhaptoglobinemia] (3)
	[Hypohaptogloginemia] (3)
141750	Alpha-thalassemia/mental retardation syndrome, type 1 (1)
141800	Erythremias, alpha- (3)
	Heinz body anemias, alpha- (3)
	Methemoglobinemias, alpha- (3)
1115==	Thalassemias, alpha- (3)
141850	Erythrocytosis (3)
	Heinz body anemia (3)
	Hemoglobin H disease (3)
L	Hypochromic microcytic anemia (3)

	Thalassemia, alpha- (3)
141900	Erythremias, beta- (3)
141900	HPFH, deletion type (3)
	Heinz body anemias, beta- (3)
	Methemoglobinemias, beta-(3)
	Sickle cell anemia (3)
	Thalassemias, beta- (3)
142000	Thalassemia due to Hb Lepore (3)
142000	Thalassemia, delta- (3)
142200	HPFH, nondeletion type A (3)
142250	HPFH, nondeletion type G (3)
142270	Hereditary persistence of fetal hemoglobin (3) (?)
142335	Hereditary persistence of fetal hemoglobin, heterocellular, Indian type (2)
142333	(?)
142380	Hepatocellular carcinoma (3)
142410	Insulin-dependent diabetes mellitus (3)
142410	MODY, type 3, 600496 (3)
	Non-insulin-dependent diabetes mellitus-2, 601407 (2)
142470	[Hereditary persistence of fetal hemoglobin, heterocellular] (2)
142600	Hemolytic anemia due to hexokinase deficiency (3)
142680	Periodic fever, familial (2)
142857	Pemphigoid, susceptibility to (2)
142858	Beryllium disease, chronic, susceptibility to (3)
142959	Hand-foot-uterus syndrome, 140000 (3)
142989	Synpolydactyly, type II, 186000 (3)
143100	Huntington disease (3)
143200	Erosive vitreoretinopathy (2)
1 13200	Wagner syndrome (2)
143450	Trifunctional protein deficiency, type II (3)
143890	Hypercholesterolemia, familial (3)
144120	Hyperimmunoglobulin G1 syndrome (2) (?)
144200	Epidermolytic palmoplantar keratoderma (3)
144700	Renal cell carcinoma (2)
145001	Hyperparathyroidism-jaw tumor syndrome (2)
145260	Pseudohypoaldosteronism, type II (2)
145410	Opitz G syndrome, type II (2)
145505	?Hypertension, essential (1)
145981	Hypocalciuric hypercalcemia, type II (2)
146150	Hypomelanosis of Ito (2) (?)
146200	Hypoparathyroidism, familial (2)
146740	Neutropenia, alloimmune neonatal (3)
	Viral infections, recurrent (3)
	Lupus erythematosus, systemic, susceptibility, 152700 (1)
146760	[IgG receptor I, phagocytic, familial deficiency of] (1)
146790	Lupus nephritis, susceptibility to (3)
147020	Agammaglobulinemia, 601495 (3)
147050	Atopy (2)
147061	Allergy and asthma susceptibility (2) (?)
147110	IgG2 deficiency, selective (3)
147141	Leukemia, acute lymphoblastic (1)
	1 ==, we are .J.mp., ve (.)

147200	[Kanna light ahain definion and (2)
147200	[Kappa light chain deficiency] (3)
147440	Growth retardation with deafness and mental retardation (3)
147450	Amytrophic lateral sclerosis, due to SOD1 deficiency, 105400 (3)
147545	Diabetes mellitus, noninsulin-dependent (3)
147557	Epidermolysis bullosa, junctional, with pyloric atresia, 226730 (3)
147570	Interferon, immune, deficiency (1)
147575	Macrocytic anemia refractory, of 5q- syndrome, 153550 (3)
	Myelodysplastic syndrome, preleukemic (3)
	Myelogenous leukemia, acute (3)
147660	Interferon, alpha, deficiency (1)
147670	Diabetes mellitus, insulin-resistant, with acanthosis nigricans (3)
	Leprechaunism (3)
	Rabson-Mendenhall syndrome (3)
147680	Severe combined immunodeficiency due to IL2 deficiency (1)
147781	Atopy, susceptibility to (3)
147790	Leukemia, acute lymphocytic, with 4/11 translocation (3) (?)
147791	Jacobsen syndrome (2)
148040	Epidermolysis bullosa simplex, Koebner, Dowling-Meara, and Weber-
	Cockayne types, 131900, 131760, 131800 (3)
148041	Pachyonychia congenita, Jadassohn-Lewandowsky type, 167200 (3)
148043	Meesmann corneal dystrophy, 122100 (3)
148065	White sponge nevus, 193900 (3)
148066	Epidermolysis bullosa simplex, Koebner, Dowling-Meara, and Weber-
	Cockayne types, 131900, 131760, 131800 (3)
	Epidermolysis bullosa simplex, recessive, 601001 (3)
148067	Nonepidermolytic palmoplantar keratoderma, 600962 (3)
	Pachyonychia congenita, Jadassohn-Lewandowsky type, 167200 (3)
148069	Pachyonychia congenita, Jackson-Lawler type, 167210 (3)
148070	?Liver disease, susceptibility to, from hepatotoxins or viruses (1)
148080	Epidermolytic hyperkeratosis, 113800 (3)
148370	Keratolytic winter erythema (2)
148500	Tylosis with esophageal cancer (2)
148900	Klippel-Feil syndrome with laryngeal malformation (2)
150000	Exertional myoglobinuria due to deficiency of LDH-A (3)
150100	Lactate dehydrogenase-B deficiency (3)
150200	[Placental lactogen deficiency] (1)
150210	Lactoferrin-deficient neutrophils, 245480 (1) (?)
150230	Langer-Giedion syndrome (2)
150240	Cutis laxa, marfanoid neonatal type (1) (?)
150250	Larsen syndrome, autosomal dominant (2)
150270	Laryngeal adductor paralysis (2) (?)
150292	Epidermolysis bullosa, Herlitz junctional type, 226700 (3)
150310	Epidermolysis bullosa, Herlitz junctional type, 226700 (3)
150510	Epidermolysis bullosa, generalized atrophic benign, 226650 (3)
151385	Leukemia, acute myeloid (3)
151383	Leukemia, acute T-cell (2)
151400	Leukemia/lymphoma, B-cell, 1 (2)
151430	Leukemia/lymphoma, B-cell, 2 (2) Leukemia, T-cell acute lymphoblastoid (2)
151440	Leukenna, 1-cen acute tymphobiastolu (2)

151670	Hepatic lipase deficiency (3)
151670	
152200	Coronary artery disease, susceptibility to (1)  Long QT syndrome-2 (3)
152427	
152445	Erythrokeratoderma, progressive symmetric, 602036 (3)
1527(0	Vohwinkel syndrome, 124500 (3)
152760	Hypogonadotropic hypogonadism due to GNRH deficiency, 227200 (1) (?)
152780	Hypogonadism, hypergonadotropic (3)  Mala manufakarmanhan ditiam dua ta defeativa I II (1) (2)
152700	Male pseudohermaphroditism due to defective LH (1) (?) Leydig cell hypoplasia (3)
152790	Precocious puberty, male, 176410 (3)
153455	Cutis laxa, recessive, type I, 219100 (1)
153700	Macular dystrophy, vitelliform type (3)
153880	Macular dystrophy, vitermorni type (3)  Macular dystrophy, dominant cystoid (2)
153900	Stargardt disease-2 (2)
154275	Malignant hyperthermia susceptibility 2 (2)
	Malignant hyperthermia susceptibility 2 (2)  Malignant hyperthermia susceptibility 3 (2)
154276	
154500	Treacher Collins mandibulofacial dysostosis (3)  Chronic infections, due to opsonin defect (3)
154545	
154550	Carbohydrate-deficient glycoprotein syndrome, type Ib, 602579 (3)  Marfan syndrome, type II (2)
154705	
155555	[Red hair/fair skin] (3)
155600	UV-induced skin damage, vulnerability to (3)  Malignant melanoma, cutaneous (2)
155600	
155900	Melkersson-Rosenthal syndrome (2) (?)
156225	Muscular dystrophy, congenital merosin-deficient (3)
156232	Mesomelic dysplasia, Kantaputra type (2)
156490	Neuroblastoma (3)  Methylcobalamin deficiency, cbl G type (3)
156570	
156600	Microcoria, congenital (2) Tietz syndrome, 103500 (3)
156845	Waardenburg syndrome, type IIA, 193510 (3)
	Waardenburg syndrome/ocular albinism, digenic, 103470 (3)
156850	Cataract, congenital, with microphthalmia (2)
157140	Dementia, frontotemporal, with parkinsonism, 601630 (3)
157147	Abetalipoproteinemia, 200100 (3)
157170	Holoprosencephaly-2 (2)
157640	PEO with mitochondrial DNA deletions, type 1 (2)
157655	Lactic acidosis due to defect in iron-sulfur cluster of complex I (1)
157900	Moebius syndrome (2) (?)
158590	Spinal muscular atrophy-4 (2)
158900	Facioscapulohumeral muscular dystrophy-1A (2)
159000	Muscular dystrophy, limb-girdle, type 1A (2)
159001	Muscular dystrophy, limb-girdle, type 1B (2)
159350	Colorectal cancer (3)
159440	Charcot-Marie-Tooth neuropathy-1B, 118200 (3)
133440	Dejerine-Sottas disease, myelin P(0)-related, 145900 (3)
	Hypomyelination, congenital (3)
159555	Leukemia, myeloid/lymphoid or mixed-lineage (2)
159595	Leukemia, transient, of Down syndrome (2)
137373	Dealerma, transferre, or Down Syllatottic (2)

160760	Cardiomyopathy, familial hypertrophic, 1, 192600 (3)
160760	
1/0777	Central core disease, one form (3) (?) Griscelli disease, 214450 (3)
160777	
160781	Cardiomyopathy, hypertrophic, mid-left ventricular chamber type (3)
160900	Myotonic dystrophy (3)
160980	Carney myxoma-endocrine complex (2)
161015	Mitochondrial complex I deficiency, 252010 (1) (?)
162100	Neuralgic amyotrophy with predilection for brachial plexus (2)
162150	Obestiy with impaired prohormone processing, 600955 (3)
162400	Neuropathy, hereditary sensory and autonomic, type 1 (2)
163729	Hypertension, pregnancy-induced (2)
163890	Parkinson disease, type 1, 601508 (3)
163950	Cardiofaciocutaneous syndrome, 115150 (2)
	Noonan syndrome-1 (2)
164009	Leukemia, acute promyelocytic, NUMA/RARA type (3)
164040	Leukemia, acute promyelocytic, NPM/RARA type (3)
164050	Nucleoside phosphorylase deficiency, immunodeficiency due to (3)
164160	Obesity, severe, due to leptin deficiency (3)
164200	Oculodentodigital dysplasia (2)
	Syndactyly, type III, 186100 (2)
164500	Spinocerebellar ataxia-7 (3)
164731	Ovarian carcinoma, 167000 (2)
164761	Hirschsprung disease, 142623 (3)
	Medullary thyroid carcinoma, 155240 (3)
	Multiple endocrine neoplasia IIA, 171400 (3)
	Multiple endocrine neoplasia IIB, 162300 (3)
164770	Myeloid malignancy, predisposition to (3)
164790	Colorectal cancer (3)
164860	Renal cell carcinoma, papillary, familial and sporadic (3)
164920	Mast cell leukemia (3)
	Mastocytosis with associated hematologic disorder (3)
	Piebaldism (3)
164953	Liposarcoma (1)
165215	3q21q26 syndrome (1)
165240	Greig cephalopolysyndactyly syndrome, 175700 (3)
	Pallister-Hall syndrome, 146510 (3)
	Postaxial polydactyly type A1, 174200 (3)
165320	Hepatocellular carcinoma (1) (?)
166600	Osteopetrosis, AD, type II (2)
166800	Otosclerosis (2)
167000	Ovarian cancer, serous (2)
167250	Paget disease of bone (2) (?)
167409	Optic nerve coloboma with renal disease, 120330 (3)
167415	Hypothyroidism, congenital, due to thyroid dysgenesis or hypoplasia (3)
168000	Paraganglioma, familial nonchromaffin, 1 (2)
168360	Paraneoplastic sensory neuropathy (1)
168450	Hypoparathyroidism, autosomal dominant(3)
100430	Hypoparathyroidism, autosomal commanu(3)
168461	Centrocytic lymphoma (2)
100401	
	Multiple myeloma, 254250 (2)

	Parathyroid adenomatosis 1 (2)
168468	Metaphyseal chondrodysplasia, Murk Jansen type, 156400 (3)
168470	Humoral hypercalcemia of malignancy (1) (?)
168500	Parietal foramina (2)
168610	Parkinsonism-dementia with pallidopontonigral degeneration (2)
169600	Hailey-Hailey disease (2)
170261	Bare lymphocyte syndrome, type I, due to TAP2 deficiency (1)
170500	Hyperkalemic periodic paralysis (3)
	Myotonia congenita, atypical acetazolamide-responsive (3)
170650	Paramyotonia congenita, 168300 (3)
170650	Periodontitis, juvenile (2)
170993	Zellweger syndrome-3 (3)
170995	Zellweger syndrome-2 (3)
171050	Colchicine resistance (3)
171060	Cholestasis, progressive familial intrahepatic, type III, 602347 (3)
171190	Hypertension, essential, 145500 (1) (?)
171650	Lysosomal acid phosphatase deficiency (1) (?)
171760	Hypophosphatasia, adult, 146300 (1) (?)
	Hypophosphatasia, infantile, 241500 (3)
171860	Hemolytic anemia due to phosphofructokinase deficiency (1)
172400	Hemolytic anemia due to glucosephosphate isomerase deficiency (3)
	Hydrops fetalis, one form (1)
172411	?Colorectal cancer, resistance to (1)
172471	Glycogenosis, hepatic, autosomal (3)
172490	Phosphorylase kinase deficiency of liver and muscle, 261750 (2) (?)
173360	Hemorrhagic diathesis due to PAI1 deficiency (1)
	Thrombophilia due to excessive plasminogen activator inhibitor (1)
173370	Plasminogen activator deficiency (1)
173470	Glanzmann thrombasthenia, type B (3)
173510	Platelet glycoprotein IV deficiency (3)
	[Macrothrombocytopenia] (1)
173610	Platelet alpha/delta storage pool deficiency (1)
173850	Polio, susceptibility to (2)
173870	Fanconi anemia (1) (?)
	Xeroderma pigmentosum (1) (?)
173910	Polycystic kidney disease, adult, type II (3)
174000	Medullary cystic kidney disease, AD (2)
174810	Osteolysis, familial expansile (2)
174900	Polyposis, juvenile intestinal (2)
175100	Adenomatous polyposis coli (3)
	Adenomatous polyposis coli, attenuated (3)
	Colorectal cancer (3)
	Desmoid disease, hereditary, 135290 (3)
	Gardner syndrome (3)
	Turcot syndrome, 276300 (3)
176000	Porphyria, acute intermittent (3)
176010	Porphyria, Chester type (2)
176100	Porphyria cutanea tarda (3)
	Porphyria, hepatoerythropoietic (3)
176260	Episodic ataxia/myokymia syndrome, 160120 (3)

176261	Jervell and Lange-Nielsen syndrome, 220400 (3)
176300	Amyloid neuropathy, familial, several allelic types (3)
1,0200	Amyloidosis, senile systemic (3)
	Carpal tunnel syndrome, familial (3)
	[Dystransthyretinemic hyperthyroxinemia](3)
176310	Leukemia, acute pre-B-cell (2)
176450	Sacral agenesis-1 (2)
176640	Creutzfeldt-Jakob disease, 123400 (3)
	Gerstmann-Straussler disease, 137440 (3)
	Insomnia, fatal familial (3)
176705	Breast cancer, sporadic (3)
176730	Diabetes mellitus, rare form (1)
	Hyperproinsulinemia, familial (3)
	MODY, one form (3)
176797	Leukemia, acute promyelocytic, PL2F/RARA type (3)
176860	Purpura fulminans, neonatal (1)
	Thrombophilia due to protein C deficiency (3)
176930	Dysprothrombinemia (3)
	Hypoprothrombinemia (3)
176943	Apert syndrome, 101200 (3)
	Beare-Stevenson cutis gyrata syndrome, 123790 (3)
	Crouzon craniofacial dysostosis, 123500 (3)
	Jackson-Weiss syndrome, 123150 (3)
1.50.1.	Pfeiffer syndrome, 101600 (3)
176947	Selective T-cell defect (3)
176960	Pituitary tumor, invasive (3)
177000	Protoporphyria, erythropoietic (3)
177070	Protoporphyria, erythropoietic, recessive, with liver failure (3)
177070	Hermansky-Pudlak syndrome, 203300 (1) (?)
177000	Spherocytosis, hereditary, Japanese type (3)
177900	Psoriasis susceptibility-1 (2)
178300	Prince of the property of the prince of the
178600	Pulmonary hypertension, familial primary (2)
178640	Pulmonary alveolar proteinosis, congenital, 265120 (3)
179095	Male infertility (1) (?)
179450 179605	Ragweed sensitivity (2) (?)  Butterfly dystrophy, retinal (3)
179003	Macular dystrophy (3)
	Retinitis pigmentosa, digenic (3)
	Retinitis pigmentosa-7, peripherin-related (3)
	Retinitis punctata albescens (3)
179615	Reticulosis, familial histocytic, 267700 (3)
	Severe combined immunodeficiency, B cell-negative, 601457 (3)
179616	Severe combined immunodeficiency, B cell-negative, 601457 (3)
179755	Renal cell carcinoma, papillary, 1 (2)
179820	[Hyperproreninemia] (3)
180069	Leber congenital amaurosis-2, 204100 (3)
10000	Retinal dystrophy, autosomal recessive, childhood-onset (3)
	Retinitis pigmentosa-20 (3)
180071	Retinitis pigmentosa, autosomal recessive (3)
	F 9

190072	Night blindness concenital stationers, type 3 163500 (3)
180072	Night blindness, congenital stationary, type 3, 163500 (3) Retinitis pigmentosa, autosomal recessive (3)
10000	Retinitis pigmentosa, autosomal recessive (3)  Retinitis pigmentosa, autosomal recessive (3)
180090	
180100	Retinitis pigmentosa-1 (2)
180104	Retinitis pigmentosa-9 (2)
180105	Retinitis pigmentosa-10 (2)
180200	Bladder cancer, 109800 (3)
	Osteosarcoma, 259500 (2)
	Pinealoma with bilateral retinoblastoma (2)
100040	Retinoblastoma (3)
180240	Leukemia, acute promyelocytic (1)
180250	Retinol binding protein, deficiency of (1) (?)
180297	Anemia, hemolytic, Rh-null, suppressor type, 268150 (3)
180380	Night blindness, congenital stationery, rhodopsin-related (3)
	Retinitis pigmentosa, autosomal recessive (3)
100201	Retinitis pigmentosa-4, autosomal dominant (3)
180381	Oguchi disease-2, 258100 (3)
180385	Leukemia, acute T-cell (2)
180721	Retinitis pigmentosa, digenic (3)
180840	Susceptibility to IDDM (1) (?)
180860	Russell-Silver syndrome (2)
180901	Central core disease, 117000 (3)
	Malignant hyperthermia susceptibility 1, 145600 (3)
181405	Scapuloperoneal spinal muscular atrophy, New England type (2)
181430	Scapuloperoneal syndrome, myopathic type (2)
181460	Schistosoma mansoni, susceptibility/resistance to (2)
181510	Schizophrenia (2) (?)
181600	Sclerotylosis (2) (?)
182138	Anxiety-related personality traits (3)
182280	Small-cell cancer of lung (2)
182290	Smith-Magenis syndrome (2)
182380	Glucose/galactose malabsorption (3)
182381	Renal glucosuria, 253100 (1) (?)
182452	Lung cancer, small cell (3)
182500	Cataract, congenital (2) (?)
182600	Spastic paraplegia-3A (2)
182601	Spastic paraplegia-4 (3)
182860	Elliptocytosis-2 (3)
	Pyropoikilocytosis (3)
	Spherocytosis, recessive (3)
182870	Anemia, neonatal hemolytic, fatal and near-fatal (3)
	Elliptocytosis-3 (3)
	Spherocytosis-1 (3)
182900	Spherocytosis-2 (3)
183600	Split hand/foot malformation, type 1 (2)
185000	Stomatocytosis I (1) (?)
185430	Atherosclerosis, susceptibility to (3) (?)
185470	Myopathy due to succinate dehydrogenase deficiency (1) (?)
185800	Symphalangism, proximal (2)

106500	A 11 4 1 (2)
186580	Arthrocutaneouveal granulomatosis (2)
186740	Immunodeficiency due to defect in CD3-gamma (3)
186770	Leukemia, T-cell acute lymphocytic (2)
186780	CD3, zeta chain, deficiency (1)
186830	Immunodeficiency, T-cell receptor/CD3 complex (3)
186855	Leukemia-2, T-cell acute lymphoblastic (3)
186860	Leukemia/lymphoma, T-cell (2)
186880	Leukemia/lymphoma, T-cell (3)
186921	Leukemia, T-cell acute lymphoblastic (2)
186940	[CD4(+) lymphocyte deficiency] (2)
	Lupus erythematosus, susceptibility to (2)
186960	Leukemia/lymphoma, T-cell (2)
187040	Leukemia-1, T-cell acute lymphoblastic (3)
188025	Thrombocytopenia, Paris-Trousseau type (2) (?)
188040	Thrombophilia due to thrombomodulin defect (3)
188070	Bleeding disorder due to defective thromboxane A2 receptor (3)
188450	Goiter, adolescent multinodular (1)
	Goiter, nonendemic, simple (3)
	Hypothyroidism, hereditary congenital (3)
188540	Hypothyroidism, nongoitrous (3)
188550	Thyroid papillary carcinoma (1)
188826	Sorsby fundus dystrophy, 136900 (3)
189800	Preeclampsia/eclampsia (2) (?)
189980	Leukemia, chronic myeloid (3)
190000	Atransferrinemia (1)
190020	Bladder cancer, 109800 (3)
190040	Dermatofibrosarcoma protuberans (3)
	Giant-cell fibroblastoma (3)
	Meningioma, SIS-related (3)
190070	Colorectal adenoma (1)
100000	Colorectal cancer (1)
190080	Burkitt lymphoma (3)
190100	Geniospasm (2)
190182	Colon cancer (3)
100105	Colorectal cancer, familial nonpolyposis, type 6 (3)
190195	Ichthyosiform erythroderma, congenital, 242100 (3)
100100	Ichthyosis, lamellar, autosomal recessive, 242300 (3)
190198	Leukemia, T-cell acute lymphoblastic (2)
190300	Tremor, familial essential, 1 (2)
190350	Trichorhinophalangeal syndrome, type I (2)
190450	Hemolytic anemia due to triosephosphate isomerase deficiency (3)
190605	Triphalangeal thumb-polysyndactyly syndrome (2)
190685	Down syndrome (1)
190900	Colorblindness, tritan (3)
191010	Cardiomyopathy, familial hypertrophic, 3, 115196 (3)
191030	Nemaline myopathy-1, 161800 (3)
191044	Cardiomyopathy, familial hypertrophic (3)
191045	Cardiomyopathy, familial hypertrophic, 2, 115195 (3)
191092	Tuberous sclerosis-2 (3)

101100	
191100	Tuberous sclerosis-1 (3)
191170	Colorectal cancer, 114500 (3)
	Li-Fraumeni syndrome (3)
191181	Cervical carcinoma (2)
191290	Segawa syndrome, recessive (3)
191315	Insensitivity to pain, congenital, with anhidrosis, 256800 (3)
191540	[Urate oxidase deficiency] (1)
192090	Breast cancer, lobular (3)
	Endometrial carcinoma (3)
	Gastric cancer, familial, 137215 (3)
	Ovarian carcinoma (3)
192340	Diabetes insipidus, neurohypophyseal, 125700 (3)
192500	Jervell and Lange-Nielsen syndrome, 220400 (3)
	Long QT syndrome-1 (3)
192974	Glycoprotein Ia deficiency (2) (?)
	Neonatal alloimmune thrombocytopenia (2)
193235	Vitreoretinopathy, neovascular inflammatory (2)
193300	Renal cell carcinoma (3)
	von Hippel-Lindau syndrome (3)
193500	Craniofacial-deafness-hand syndrome, 122880 (3)
	Rhabdomyosarcoma, alveolar, 268220 (3)
	Waardenburg syndrome, type I (3)
	Waardenburg syndrome, type III, 148820 (3)
194070	Denys-Drash syndrome (3)
	Frasier syndrome, 136680 (3)
	Wilms tumor, type 1 (3)
194071	Adrenocortical carcinoma, hereditary, 202300 (2)
	Wilms tumor, type 2 (2)
194190	Wolf-Hirschhorn syndrome (2)
200150	Choreoacanthocytosis (2)
200350	Acetyl-CoA carboxylase deficiency (1)
200990	Acrocallosal syndrome (2) (?)
201450	Acyl-CoA dehydrogenase, medium chain, deficiency of (3)
201460	Acyl-CoA dehydrogenase, long chain, deficiency of (3)
201470	Acyl-CoA dehydrogenase, short-chain, deficiency of (3)
201475	VLCAD deficiency (3)
201810	3-beta-hydroxysteroid dehydrogenase, type II, deficiency (3)
201910	Adrenal hyperplasia, congenital, due to 21-hydroxylase deficiency (3)
202010	Adrenal hyperplasia, congenital, due to 11-beta-hydroxylase deficiency (3)
	Aldosteronism, glucocorticoid-remediable (3)
203100	Albinism, oculocutaneous, type IA (3)
	Waardenburg syndrome/ocular albinism, digenic, 103470 (3)
203310	Ocular albinism, autosomal recessive (2) (?)
203500	Alkaptonuria (3)
203740	Alpha-ketoglutarate dehydrogenase deficiency (1)
203750	3-ketothiolase deficiency (3)
203800	Alstrom syndrome (2)
204500	Ceroid-lipofuscinosis, neuronal 2, classic late infantile (2)
205100 205900	Amyotrophic lateral sclerosis, juvenile (2) Anemia, Diamond-Blackfan (2)

00	
207750	Hyperlipoproteinemia, type Ib (3)
207800	Argininemia (3)
208100	Arthrogryposis multiplex congenita, neurogenic (2)
208250	Jacobs syndrome (2)
208400	Aspartylglucosaminuria (3)
208900	Ataxia-telangiectasia (3)
	B-cell non-Hodgkin lymphoma, sporadic (3)
	T-cell prolymphocytic leukemia, sporadic (3)
209900	Bardet-Biedl syndrome 2 (2)
209901	Bardet-Biedl syndrome 1 (2)
210900	Bloom syndrome (3)
211420	Breast cancer, ductal (2)
212138	Carnitine-acylcarnitine translocase deficiency (3)
212200	Carnosinemia (2)
213700	Cerebrotendinous xanthomatosis (3)
214300	Klippel-Feil syndrome (2) (?)
214400	Charcot-Marie-Tooth neuropathy-4A (2)
214500	Chediak-Higashi syndrome (3)
215700	Citrullinemia (3)
216550	Cohen syndrome (2)
216900	Achromatopsia (2)
216950	C1r/C1s deficiency, combined (1)
217000	C2 deficiency (3)
217030	C3b inactivator deficiency (3)
217050	C6 deficiency (1)
217020	Combined C6/C7 deficiency (1)
217070	C7 deficiency (1)
217800	Macular corneal dystrophy (2)
218000	Andermann syndrome (2)
218030	Apparent mineralocorticoid excess, hypertension due to (3)
219800	Cystinosis, nephropathic (3)
221770	Polycystic lipomembranous osteodysplasia with sclerosing
221,,0	leukencephalopathy (2)
221820	Gliosis, familial progressive subcortical (2)
222100	Diabetes mellitus, insulin-dependent-1 (2) (?)
222600	Achondrogenesis Ib, 600972 (3)
	Atelosteogenesis II, 256050 (3)
	Diastrophic dysplasia (3)
222700	Lysinuric protein intolerance (2)
222745	DECR deficiency (2) (?)
222800	Hemolytic anemia due to bisphosphoglycerate mutase deficiency (1)
222900	Sucrose intolerance (3)
223000	Lactase deficiency, adult, 223100 (1) (?)
223000	Lactase deficiency, congenital (1) (?)
223360	Dopamine-beta-hydroxylase deficiency (1)
223900	Dysautonomia, familial (2)
224100	Congenital dyserythropoietic anemia II (2)
224100	Dyserythropoietic anemia, contenital, type I (2)
225500	Ellis-van Creveld syndrome (2)
223300	Ems-van Cieveid Syndrome (2)

227220	Epidermolysis bullosa inversa, junctional (2)
	fr 1 1 3 (2)
	[Eye color, brown] (2)
	Hemorrhagic diathesis due to factor V deficiency (1)
	Thromboembolism susceptibility due to factor V Leiden (3)
	Factor VII deficiency (3)
	Factor X deficiency (3)
	Fanconi anemia, type C (3)
	Fanconi anemia, type D (2)
	Fanconi anemia, type A (3)
229000	Fletcher factor deficiency (1)
	Friedreich ataxia (3)
	Friedreich ataxia with retained reflexes (2)
	Fructose intolerance (3)
229700	Fructose-bisphosphatase deficiency (1)
230000	Fucosidosis (3)
230200	Galactokinase deficiency with cataracts (3)
	Galactose epimerase deficiency (3)
230450	Hemolytic anemia due to gamma-glutamylcysteine synthetase deficiency
	(1)
	GM1-gangliosidosis (3)
	Mucopolysaccharidosis IVB (3)
	Gaucher disease (3)
	Gaucher disease with cardiovascular calcification (3)
231200	Bernard-Soulier syndrome (3)
231550	Achalasia-addisonianism-alacrimia syndrome (2)
231670	Glutaricaciduria, type I (3)
231675	Glutaricaciduria, type IIC (3)
ļ	Glutaricaciduria, type IIA (1)
	Glutathioninuria (1)
	Propionicacidemia, type I or pccA type (1)
	Propionicacidemia, type II or pccB type (3)
	Glycogen storage disease I (3)
	Glycogen storage disease IIIa (1)
	Glycogen storage disease IIIb (3)
232600	McArdle disease (3)
232700	Glycogen storage disease VI (3)
232700	Glycogen storage disease VII (3)
233100	[Renal glucosuria] (2)
233690	Chronic granulomatous disease, autosomal, due to deficiency of CYBA (3)
233700	Chronic granulomatous disease, autosomai, due to deficiency of CTBA (3)
233710	Chronic granulomatous disease due to deficiency of NCF-1 (3)  Chronic granulomatous disease due to deficiency of NCF-2 (1)
	Factor XII deficiency (3)
234000	Neurodegeneration with brain iron accumulation (2)
235200	Hemochromatosis (3)
235800	[Histidinemia] (1)
236100	Holoprosencephaly-1 (2)
236200	Homocystinuria, B6-responsive and nonresponsive types (3)
236700	McKusick-Kaufman syndrome (2)
236730	Urofacial syndrome (2)

227200	Coult are order to complete country (2)
237300	Carbamoylphosphate synthetase I deficiency (3)
238300	Hyperglycinemia, nonketotic, type I (3)
238310	Hyperglycinemia, nonketotic, type II (1)
238600	Chylomicronemia syndrome, familial (3)
	Combined hyperlipemia, familial (3)
	Hyperlipoproteinemia I (1)
220070	Lipoprotein lipase deficiency (3)
238970	HHH syndrome (2) (?)
239500	Hyperprolinemia, type I (1)
240300	Autoimmune polyglandular disease, type I (3)
243500	Isovalericacidemia (3)
245000	Papillon-Lefevre syndrome (2)
245050	Ketoacidosis due to SCOT deficiency (3)
245200	Krabbe disease (3)
245349	Lacticacidemia due to PDX1 deficiency (3)
245900	Fish-eye disease (3)
	Norum disease (3)
246450	HMG-CoA lyase deficiency (3)
246530	Leukotriene C4 synthase deficiency (1)
246900	Lipoamide dehydrogenase deficiency (3)
247200	Miller-Dieker lissencephaly syndrome (2)
247640	Leukemia, acute lymphoblastic (2)
248510	Mannosidosis, beta- (3)
248600	Maple syrup urine disease, type Ia (3)
248610	Maple syrup urine disease, type II (3)
248611	Maple syrup urine disease, type Ib (3)
249000	Meckel syndrome (2)
249270	Thiamine-responsive megaloblastic anemia (2)
250100	Metachromatic leukodystrophy (3)
250800	Methemoglobinemia, type I (3)
	Methemoglobinemia, type II (3)
250850	Hypermethioninemia, persistent, autosomal dominant, due to methionine
	adenosyltransferase I/III deficiency (3)
251000	Methylmalonicaciduria, mutase deficiency type (3)
251170	Mevalonicaciduria (3)
252500	Mucolipidosis II (1)
	Mucolipidosis III (1)
252800	Mucopolysaccharidosis Ih (3)
	Mucopolysaccharidosis Ih/s (3)
	Mucopolysaccharidosis Is (3)
252920	Sanfilippo syndrome, type B (3)
252940	Sanfilippo syndrome, type D (1)
253000	Mucopolysaccharidosis IVA (3)
253200	Maroteaux-Lamy syndrome, several forms (3)
253220	Mucopolysaccharidosis VII (3)
253250	Mulibrey nanism (2)
253260	Biotinidase deficiency (3)
253270	Multiple carboxylase deficiency, biotin-responsive (3)
253601	Miyoshi myopathy, 254130 (2)
	1 J

	Muscular dystrophy, limb-girdle, type 2B (2)
252700	Muscular dystrophy, limb-girdle, type 2C (3)
253700	
253800	Fukuyama type congenital muscular dystrophy (2)
254210	Walker-Warburg syndrome, 236670 (2) (?)
254210	Myasthenia gravis, familial infantile (2)
254770	Epilepsy, juvenile myoclonic (2)
255800	Schwartz-Jampel syndrome (2)
256030	Nemaline myopathy-2 (2)
256100	Nephronophthisis, juvenile (3)
256540	Galactosialidosis (3)
256550	Sialidosis, type I (3)
	Sialidosis, type II (3)
256731	Ceroid-lipofuscinosis, neuronal-5, variant late infantile (3)
257200	Niemann-Pick disease, type A (3)
	Niemann-Pick disease, type B (3)
257220	Niemann-Pick disease, type C.(3)
	Niemann-Pick disease, type D, 257250 (2)
258501	3-methylglutaconicaciduria, type III (2)
258870	Gyrate atrophy of choroid and retina with ornithinemia, B6 responsive or
	unresponsive (3)
258900	Oroticaciduria (3)
259700	Osteopetrosis, recessive (2)
259730	Renal tubular acidosis-osteopetrosis syndrome (3)
259770	Osteoporosis-pseudoglioma syndrome (2)
259900	Hyperoxaluria, primary, type 1 (3)
261510	Pseudo-Zellweger syndrome (1)
261600	Phenylketonuria (3)
	[Hyperphenylalaninemia, mild] (3)
261640	Phenylketonuria due to PTS deficiency (3)
261670	Myopathy due to phosphoglycerate mutase deficiency (3)
262000	Bjornstad syndrome (2)
262850	Plasmin inhibitor deficiency (3)
263200	Polycystic kidney disease, autosomal recessive (2)
263700	Porphyria, congenital erythropoietic (3)
264300	Pseudohermaphroditism, male, with gynecomastia (3)
264470	Adrenoleukodystrophy, pseudoneonatal (2)
264600	Pseudovaginal perineoscrotal hypospadias (3)
264700	Pseudo-vitamin D dependency rickets 1 (2)
264900	Factor XI deficiency (3)
266100	Pyridoxine dependency with seizures (1) (?)
266150	Pyruvate carboxylase deficiency (3)
266200	Anemia, hemolytic, due to PK deficiency (3)
266300	[Hair color, red] (2)
266600	Inflammatory bowel disease-1 (2)
267750	Knobloch syndrome (2)
268800	Sandhoff disease, infantile, juvenile, and adult forms (3)
200000	Spinal muscular atrophy, HEXB-related (3)
268900	[Sarcosinemia] (2)

270200	Singram Largeon gyndroma (2)
270200	Sjogren-Larsson syndrome (3) Spastic paraplegia-5A (2)
	Spinocerebellar ataxia-8, infantile, with sensory neuropathy (2)
271245	
271900	Canavan disease (3) GM2-gangliosidosis, AB variant (3)
272750	
272800	GM2-gangliosidosis, juvenile, adult (3)
	Tay-Sachs disease (3)
272200	[Hex A pseudodeficiency] (1)
273300	Male germ cell tumor (2) Glanzmann thrombasthenia, type A (3)
273800	Thrombocytopenia, neonatal alloimmune (1)
274180	Thromboxane synthase deficiency (2)
	Thromboxane synthase deficiency (2)  Thymine-uraciluria (1)
274270	Fluorouracil toxicity, sensitivity to (1)
274500	Goiter, congenital (3)
2/4300	Hypothyroidism, congenital (3)
	Thyroid iodine peroxidase deficiency (1)
274600	Deafness, autosomal recessive 4 (3)
274000	Pendred syndrome (3)
275200	Graves disease, 275000 (1)
273200	Hyperthroidism, congenital (3)
	Hypothyroidism, nongoitrous, due to TSH resistance (3)
	Thyroid adenoma, hyperfunctioning (3)
275350	Transcobalamin II deficiency (3)
276000	Pancreatitis, hereditary, 167800 (3)
270000	Trypsinogen deficiency (1)
276600	Tyrosinemia, type II (3)
276700	Tyrosinemia, type I (3)
276710	Tyrosinemia, type III (1)
276901	Usher syndrome, type 2 (3)
276902	Usher syndrome, type 3 (2)
276903	Deafness, autosomal dominant 11, neurosensory, 601317 (3)
270903	Deafness, autosomal recessive 2, neurosensory, 600060 (3)
	Usher syndrome, type 1B (3)
276904	Usher syndrome, type 1C (2)
277700	Werner syndrome (3)
277730	Wernicke-Korsakoff syndrome, susceptibility to (1)
277900	Wilson disease (3)
278000	Cholesteryl ester storage disease (3)
	Wolman disease (3)
278250	Wrinkly skin syndrome (2)
278300	Xanthinuria, type I (3)
278700	Xeroderma pigmentosum, group A (3)
278720	Xeroderma pigmentosum, group C (3)
278760	Xeroderma pigmentosum, group F (3)
300011	Cutis laxa, neonatal (3)
	Menkes disease, 309400 (3)
	Occipital horn syndrome, 304150 (3)
300029	Retinitis pigmentosa-15 (2)
300031	Mental retardation, X-linked, FRAXF type (3)
	1

Alpha-thalassemia/mental retardation syndrome, type 2, 301040 (3)
Juberg-Marsidi syndrome, 309590 (3)
?Wernicke-Korsakoff syndrome, susceptibility to (1)
Mental retardation, X-linked 23, nonspecific (2)
Mental retardation, X-linked 20 (2)
Intestinal pseudoobstruction, neuronal, X-linked (2)
BPNH/MR syndrome (2)
Nodular heterotopia, bilateral periventricular (2)
Mental retardation with psychosis, pyramidal signs, and macroorchidism (2)
Night blindness, congenital stationary, type 2 (2)
Coffin-Lowry syndrome, 303600 (3)
Mental retardation, X-linked 29 (2)
Cone dystrophy, progressive X-linked, 2 (2)
Epilepsy, female restricted, with mental retardation (2)
Adrenoleukodystrophy (3)
Adrenomyeloneuropathy (3)
Mental retardation, X-linked nonspecific, 309541 (3)
Night blindness, congenital stationary, X-linked incomplete, 300071 (3)
Mental retardation with isolated growth hormone deficiency (2)
Dyskeratosis congenita-1, 305000 (3)
Agammaglobulinemia, type 1, X-linked (3)
XLA and isolated growth hormone deficiency, 307200 (3) (?)
Ocular albinism, Forsius-Eriksson type (2)
Albinism-deafness syndrome (2)
Thrombocytopenia, X-linked, 313900 (3)
Wiskott-Aldrich syndrome (3)
Amelogenesis imperfecta (3)
Amelogenesis imperfecta-3, hypoplastic type (2) (?)
Anemia, sideroblastic/hypochromic (3)
Anemia, sideroblastic, with spinocerebellar ataxia (2) (?)
Fabry disease (3)
Anophthalmos-1 (2) (?)
Arthrogryposis, X-linked (spinal muscular atrophy, infantile, X-linked) (2)
Arts syndrome (2)
Bazex syndrome (2)
Borjeson-Forssman-Lehmann syndrome (2)
Barth syndrome (3)
Cardiomyopathy, X-linked dilated, 300069 (3)
Endocardial fibroelastosis-2 (2)
Noncompaction of left ventricular myocardium, isolated (3)
Nance-Horan syndrome (2)
Charcot-Marie-Tooth neuropathy, X-linked-2, recessive (2)
Chondrodysplasia punctata, X-linked dominant (2)
Cleft palate, X-linked (2)
Alport syndrome, 301050 (3)
Leiomyomatosis-nephropathy syndrome, 308940 (1)
Leiomyomatosis, diffuse, with Alport syndrome (3)

303800	Colorblindness, deutan (3)
303900	Colorblindness, protan (3)
304040	Charcot-Marie-Tooth neuropathy, X-linked-1, dominant, 302800 (3)
304340	Mental retardation, X-linked, syndromic-5, with Dandy-Walker
	malformation, basal ganglia disease, and seizures (2)
304500	Deafness, X-linked 2, perceptive congenital (2)
304700	Deafness, X-linked 1, progressive (3)
	Jensen syndrome, 311150 (3)
	Mohr-Tranebjaerg syndrome (3)
304800	Diabetes insipidus, nephrogenic (3)
305100	Anhidrotic ectodermal dysplasia (2)
305400	Aarskog-Scott syndrome (3)
305435	Heterocellular hereditary persistence of fetal hemoglobin, Swiss type (2)
305450	FG syndrome (2)
305900	Favism (3)
	G6PD deficiency (3)
	Hemolytic anemia due to G6PD deficiency (3)
306000	Glycogenosis, X-linked hepatic, type I (3)
	Glycogenosis, X-linked hepatic, type II (3)
306100	Gonadal dysgenesis, XY female type (2)
306250	Leukemia, acute myeloid, M2 type (1)
306700	Hemophilia A (3)
306900	Hemophilia B (3)
306955	Heterotaxy, X-linked visceral (3)
306995	[?Homosexuality, male] (2)
307150	Hypertrichosis, congenital generalized (2)
307700	Hypoparathyroidism, X-linked (2)
307800	Hypophosphatemia, hereditary (3)
308000	HPRT-related gout (3)
	Lesch-Nyhan syndrome (3)
308100	Ichthyosis, X-linked (3)
	Placental steroid sulfatase deficiency (3)
308240	Lymphoproliferative syndrome, X-linked (2)
308300	Incontinentia pigmenti, sporadic type (2)
308310	Incontinentia pigmenti, familial (2)
308380	Combined immunodeficiency, X-linked, moderate, 312863 (3)
	Severe combined immunodeficiency, X-linked, 300400 (3)
308800	Keratosis follicularis spinulosa decalvans (2)
308840	Hydrocephalus due to aqueductal stenosis, 307000 (3)
	MASA syndrome, 303350 (3)
200000	Spastic paraplegia, 312900 (3)
309000	Lowe syndrome (3)  Mario demossive illness V linked (2) (2)
309200	Manic-depressive illness, X-linked (2) (?)
309300	Megalocornea, X-linked (2)
309470	Mental retardation, X-linked, syndromic-3, with spastic diplegia (2)
309500	Renpenning syndrome-1 (2)
309510	Mental retardation, X-linked, syndromic-1, with dystonic movements,
200540	ataxia, and seizures (2)
309548	Mental retardation, X-linked, FRAXE type (3)
309605	Mental retardation, X-linked, syndromic-4, with congenital contractures

	and low fingertip arches (2)
309610	Mental retardation, X-linked, syndromic-2, with dysmorphism and cerebral
307010	atrophy (2)
309620	Mental retardation-skeletal dysplasia (2)
309850	Brunner syndrome (3)
309900	Mucopolysaccharidosis II (3)
310300	Emery-Dreifuss muscular dystrophy (3)
310400	Myotubular myopathy, X-linked (3)
310460	Bornholm eye disease (2)
310400	Myopia-1 (2)
310490	Cowchock syndrome (2)
311050	Optic atrophy, X-linked (2)
311200	Oral-facial-digital syndrome 1 (2)
311300	Otopalatodigital syndrome, type I (2)
311510	Waisman parkinsonism-mental retardation syndrome (2)
311770	Paroxysmal nocturnal hemoglobinuria (3)
311800	Hemolytic anemia due to PGK deficiency (3)
311000	Myoglobinuria/hemolysis due to PGK deficiency (3)
311850	Phosphoribosyl pyrophosphate synthetase-related gout (3)
311870	Muscle glycogenosis (3)
312000	Panhypopituitarism, X-linked (2)
312040	N syndrome, 310465 (1) (?)
312060	Properdin deficiency, X-linked (3)
312080	Pelizaeus-Merzbacher disease (3)
312000	Spastic paraplegia-2, 312920 (3)
312170	Pyruvate dehydrogenase deficiency (3)
312700	Retinoschisis (3)
312760	Turner syndrome (1)
312865	Langer mesomelic dysplasia, 249700 (3)
312003	Leri-Weill dyschondrosteosis, 127300 (3)
	Short stature, idiopathic familial (3)
313400	Spondyloepiphyseal dysplasia tarda (2)
313850	Thoracoabdominal syndrome (2)
314250	Dystonia-3, torsion, with parkinsonism, Filipino type (2)
314300	Goeminne TKCR syndrome (2)
314400	Cardiac valvular dysplasia-1 (2)
314580	Wieacker-Wolff syndrome (2)
600020	Prostate cancer, 176807 (3)
600040	Colorectal cancer (3)
600045	Xeroderma pigmentosum, group E, subtype 2 (1)
600048	Breast cancer-3 (2)
600049	Myelodysplasia syndrome-1 (3)
600059	Retinitis pigmentosa-13 (2)
600065	Leukocyte adhesion deficiency, 116920 (3)
600079	Colon cancer (3)
600095	Split hand/foot malformation, type 3 (2)
600101	Deafness, autosomal dominant 2 (2)
600105	Retinitis pigmentosa-12, autosomal recessive (2)
600119	Adhalinopathy, primary (1)
000117	

	Muscular dystrophy, Duchenne-like, type 2 (3)
600138	Retinitis pigmentosa-11 (2)
600140	Rubenstein-Taybi syndrome, 180849 (3)
600143	Epilepsy, progressive, with mental retardation (2)
600160	Melanoma, 155601 (3)
600163	Long QT syndrome-3 (3)
600173	SCID, autosomal recessive, T-negative/B-positive type (3)
600175	Spinal muscular atrophy, congenital nonprogressive, of lower limbs (2)
600179	Leber congenital amaurosis, type I, 204000 (3)
600184	Carnitine acetyltransferase deficiency (1) (?)
600185	Breast cancer 2, early onset (3)
000103	Pancreatic cancer (3)
600192	Sarcoma, synovial (1)
600194	Ichthyosis bullosa of Siemens, 146800 (3)
600202	Dyslexia, specific, 2 (2)
600211	Cleidocranial dysplasia, 119600 (3)
600221	Venous malformations, multiple cutaneous and mucosal, 600195 (3)
600223	Spinocerebellar ataxia-4 (2)
600228	Pseudohypoaldosteronism, type I, 264350 (3)
600231	Palmoplantar keratoderma, Bothnia type (2)
600234	HMG-CoA synthease-2 deficiency (1)
600243	Temperature-sensitive apoptosis (1)
600258	Colorectal cancer, hereditary nonpolyposis, type 3 (3)
600261	Ehlers-Danlos-like syndrome (3)
600266	Resistance/susceptibility to TB, etc. (1) (?)
600273	Polycystic kidney disease, infantile severe, with tuberous sclerosis (3)
600276	Cerebral arteriopathy with subcortical infarcts and leukoencephalopathy,
000270	125310 (3)
600281	MODY, type 1, 125850 (3)
	Non-insulin-dependent diabetes mellitus, 125853 (3)
600309	Atrioventricular canal defect-1 (2)
600310	Epiphyseal dysplasia, multiple 1, 132400 (3)
	Pseudoachondroplasia, 177170 (3)
600318	Diabetes mellitus, insulin-dependent, 3 (2)
600319	Diabetes mellitus, insulin-dependent, 4 (2)
600320	Insulin-dependent diabetes mellitus-5 (2)
600321	Diabetes mellitus, insulin-dependent, 7 (2)
600332	Rippling muscle disease-1 (2)
600354	Spinal muscular atrophy-1, 253300 (3)
	Spinal muscular atrophy-2, 253550 (3)
	Spinal muscular atrophy-3, 253400 (3)
600359	Bartter syndrome, type 2 (3)
600364	Cone dystrophy-3, 602093 (3)
600374	Bardet-Biedl syndrome 4 (2)
600414	Adrenoleukodystrophy, neonatal, 202370 (3)
600415	Ataxia with isolated vitamin E deficiency, 277460 (3)
600429	[li blood group, 110800] (1)
600430	Brachydactyly-mental retardation syndrome (2)
600467	Malignant hyperthermia susceptibility 4 (2)

500500	TD 11 11 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1
600509	Persistent hyperinsulinemic hypoglycemia of infancy, 256450 (3)
600510	Pigment dispersion syndrome (2)
600512	Epilepsy, partial (2)
600525	Trichodontoosseous syndrome, 190320 (3)
600528	CPT deficiency, hepatic, type I, 255120 (1)
600536	Myopathy, congenital (3)
600542	Chondrosarcoma, extraskeletal myxoid (1)
600584	Atrial septal defect with atrioventricular conduction defects, 108900 (3)
600593	Craniosynostosis, Adelaide type (2)
600617	Lipoid adrenal hyperplasia, 201710 (3)
600618	Leukemia, acute lymphoblastic (1)
600623	Prostate cancer, 176807 (2)
600624	Cone-rod retinal dystrophy-1 (2)
600631	Enuresis, nocturnal, 1 (2)
600635	Goiter, familial, due to TTF-1 defect (1)
600650	CPT deficiency, hepatic, type II, 600649 (3)
	Myopathy due to CPT II deficiency, 255110 (3)
600652	Deafness, autosomal dominant 4 (2)
600678	Cancer susceptibility (3)
600698	Lipoma (3)
	Lipomatosis, mutiple, 151900 (2) (?)
	Salivary adenoma (3)
	Uterine leiomyoma (3)
600701	Lipoma (1) (?)
600722	Ceroid lipofuscinosis, neuronal, variant juvenile type, with granular
	osmiophilic deposits (3)
	Ceroid lipofuscinosis, neuronal-1, infantile, 256730 (3)
600725	Holoprosencephaly-3, 142945 (3)
600757	Orofacial cleft-3 (2)
600759	Alzheimer disease-4 (3)
600760	Liddle syndrome, 177200 (3)
	Pseudohypoaldosteronism, type I, 264350 (3)
600761	Liddle syndrome, 177200 (3)
	Pseudohypoaldosteronism, type I, 264350 (3)
600792	Deafness, autosomal recessive 5 (2)
600805	Epidermolysis bullosa, junctional, Herlitz type (3)
600807	Bronchial asthma (2)
600808	Enuresis, nocturnal, 2 (2)
600811	Xeroderma pigmentosum, group E, DDB-negative subtype, 278740 (3)
600835	AIDS, resistance to (3)
600837	Hirschsprung disease, 142623 (3)
600839	Bartter syndrome, 241200 (3)
600850	Schizophrenia disorder-4 (2)
600852	Retinitis pigmentosa-17 (2)
600856	Beckwith-Wiedemann syndrome, 130650 (3)
600857	Leigh syndrome (3)
600881	Cataract, congenital, zonular, with sutural opacities (2)
600882	Charcot-Marie-Tooth neuropathy-2B (2)
600883	Diabetes mellitus, insulin-dependent, 8 (2)

600004	Cardiomyopathy, familial dilated 1B (2)
600884	
600887	Endometrial carcinoma (3)
600890	LCHAD deficiency (3)
600007	Mitochondrial trifunctional protein deficiency (1)
600897	Cataract, zonular pulverulent-1, 116200 (3)
600899	Severe combined immunodeficiency, type I, 202500 (1) (?)
600900	Muscular dystrophy, limb-girdle, type 2E (3)
600918	Cystinuria, type III (2)
600919	Long QT syndrome-4 with sinus bradycardia (2)
600923	Porphyria variegata, 176200 (3)
600937	Persistent hyperinsulinemic hypoglycemia of infancy, 256450 (3)
600946	Laron dwarfism, 262500 (3)
	Short stature, autosomal dominant, with normal serum growth hormone
	binding protein (3)
	Short stature, idiopathic (3)
600956	Persistent Mullerian duct syndrome, type II, 261550 (3)
600957	Persistent Mullerian duct syndrome, type I, 261550 (3)
600958	Cardiomyopathy, familial hypertrophic, 4, 115197 (3)
600964	Refsum disease, adult, with increased pipecolicacidemia (2)
600965	Deafness, autosomal dominant 6 (2)
600968	Gitelman syndrome, 263800 (3)
600971	Deafness, autosomal recessive 6 (2)
600974	Deafness, autosomal recessive 7 (2)
600975	Glaucoma 3, primary infantile, B (2)
600977	Cone dystrophy, progressive (2)
600983	Pseudohypoaldosteronism type I, autosomal dominant, 177735 (3)
600993	Pancreatic cancer (3)
600994	Deafness, autosomal dominant 5 (2)
600995	Nephrotic syndrome, idiopathic, steroid-resistant (2)
600996	Arrhythmogenic right ventricular dysplasia-2 (2)
600998	Bleeding diathesis due to GNAQ deficiency (1)
601002	5-oxoprolinuria, 266130 (3)
001002	Hemolytic anemia due to glutathione synthetase deficiency, 231900 (3)
601011	Cerebellar ataxia, pure (3)
00.011	Episodic ataxia, type 2, 108500 (3)
	Hemiplegic migraine, familial, 141500 (3)
	Spinocerebellar ataxia-6, 183086 (3)
601071	Deafness, autosomal recessive 9 (2)
601072	Deafness, autosomal recessive 8 (2)
601097	Charcot-Marie-Tooth neuropathy-1A, 118220 (3)
	Dejerine-Sottas disease, PMP22 related, 145900 (3)
	Neuropathy, recurrent, with pressure palsies, 162500 (3)
601105	Pycnodysostosis, 265800 (3)
601107	Dubin-Johnson syndrome, 237500 (3)
601130	Tolbutamide poor metabolizer (3)
601145	Epilepsy, progressive myoclonic 1, 254800 (3)
601146	Acromesomelic dysplasia, Hunter-Thompson type, 201250 (3)
222.0	Brachydactyly, type C, 113100 (3)
	Chondrodysplasia, Grebe type, 200700 (3)
601154	Cardiomyopathy, dilated, 1E (2)

601199	Hypocalcemia, autosomal dominant, 601198 (3)
	Hypocalciuric hypercalcemia, type I, 145980 (3)
	Neonatal hyperparathyroidism, 239200 (3)
601202	Cataract, anterior polar-2 (2)
601208	Insulin-dependent diabetes mellitus-11 (2)
601226	Progressive external ophthalmoplegia, type 2 (2)
601238	Cerebellar ataxia, Cayman type (2)
601253	Muscular dystrophy, limb-girdle, type IC (3)
601267	HIV infection, susceptibility/resistence to (3)
601277	Ichthyosis, lamellar, type 2 (2)
601284	Hereditary hemorrhagic telangiectasia-2, 600376 (3)
601295	Bile acid malabsorption, primary (3)
601309	Basal cell carcinoma, sporadic (3)
	Basal cell nevus syndrome, 109400 (3)
601313	Polycystic kidney disease, adult type I, 173900 (3)
601316	Deafness, autosomal dominant 10 (2)
601318	Diabetes mellitus, insulin-dependent, 13 (2)
601362	DiGeorge syndrome/velocardiofacial syndrome complex-2 (2)
601363	Wilms tumor, type 4 (2)
601369	Deafness, autosomal dominant 9 (2)
601373	HIV infection, susceptibility/resistance to (3)
601382	Charcot-Marie-Tooth neuropathy-4B (2)
601385	Prostate cancer (1) (?)
601386	Deafness, autosomal recessive 12 (2)
601387	Breast cancer (3)
601399	Platelet disorder, familial, with associated myeloid malignancy (2)
601406	B-cell non-Hodgkin lymphoma, high-grade (3)
601410	Diabetes mellitus, transient neonatal (2)
601411	Muscular dystrophy, limb-girdle, type 2F, 601287 (3)
601412	Deafness, autosomal dominant 7 (2)
601414	Retinitis pigmentosa-18 (2)
601471	Moebius syndrome-2 (2)
601472	Charcot-Marie-Tooth neuropathy-2D (2)
601493	Cardiomyopathy, dilated 1C (2)
601494	Cardiomyopathy, familial, dilated-2 (2)
601498	Peroxisomal biogenesis disorder, complementation group 4 (3)
601499	Rieger syndrome, type 2 (2)
601517	Spinocerebellar ataxia-2, 183090 (3)
601518	Prostate cancer, hereditary, 1, 176807 (2)
601542	Rieger syndrome, type 1, 180500 (3)
601545	Lissencephaly-1 (3)
601567	Combined factor V and VIII deficiency, 227300 (3)
601596	Charcot-Marie-Tooth neuropathy, demyelinating (2)
601604	Mycobacterial and salmonella infections, susceptibility to (3)
601606	Trichoepithelioma, multiple familial (2)
601620	Holt-Oram syndrome, 142900 (3)
601621	Ulnar-mammary syndrome, 181450 (3)
001021	Omar-maintary syndronic, 101750 (5)
601622	Saethre-Chotzen syndrome, 101400 (3)

WO 01/22920 PCT/US00/26524

403

601649	Blepharophimosis, epicanthus inversus, and ptosis, type 2 (2)
601650	Paraganglioma, familial nonchromaffin, 2 (2)
601652	Glaucoma 1A, primary open angle, juvenile-onset, 137750 (3)
601653	Branchiootic syndrome (3)
	Branchiootorenal syndrome, 113650 (3)
601666	Insulin-dependent diabetes mellitus-15 (2)
601669	Hirschsprung disease, one form (2) (?)
601676	Acute insulin response (2)
601680	Distal arthrogryposis, type 2B (2)
601682	Glaucoma 1C, primary open angle (2)
601687	Meesmann corneal dystrophy, 122100 (3)
601690	Platelet-activating factor acetylhydrolase deficiency (3)
601691	Cone-rod dystrophy 3 (3)
	Fundus flavimaculatus with macular dystrophy, 248200 (3)
	Retinitis pigmentosa-19, 601718 (3)
	Stargardt disease-1, 248200 (3)
601692	Corneal dystrophy, Avellino type (3)
	Corneal dystrophy, Groenouw type I, 121900 (3)
	Corneal dystrophy, lattice type I, 122200 (3)
	Reis-Bucklers corneal dystrophy (3)
601718	Retinitis pigmentosa-19 (2)
601744	Systemic lupus erythematosus, susceptibility to, 1 (2)
601757	Rhizomelic chondrodysplasia punctata, type 1, 215100 (3)
601768	Leukemia, acute myeloid (3)
601769	Osteoporosis, involutional (1) (?)
	Rickets, vitamin D-resistant, 277440 (3)
601771	Glaucoma 3A, primary infantile, 231300 (3)
601777	Cone dystrophy, progressive (2)
601780	Ceroid-lipofuscinosis, neuronal-6, variant late infantile (2)
601785	Carbohydrate-deficient glycoprotein syndrome, type I, 212065 (3)
601800	[Hair color, brown] (2)
601841	Protein C inhibitor deficiency (2)
601843	Hypothyroidism, congenital, 274400 (3)
601844	Pseudohypoaldosteronism type II (2)
601846	Muscular dystrophy with rimmed vacuoles (2)
601847	Progressive intrahepatic cholestasis-2 (2)
601850	Retinitis pigmentosa-deafness syndrome (2)
601863	Bare lymphocyte syndrome, complementation group C (1)
601868	Deafness, autosomal dominant 13 (2)
601884	[High bone mass] (2)
601885	Cataract, zonular pulverulent-2 (2)
601889	Lymphoma, diffuse large cell (3)
601916	Pancreatic cancer (2)
601920	Alagille syndrome, 118450 (3)
601928	Monilethrix, 158000 (3)
601941	Insulin-dependent diabetes mellitus-6 (2)
601954	Muscular dystrophy, limb-girdle, type 2G (2)
601969	Glioblastoma multiforme, 137800 (3)
	Medulloblastoma, 155255 (3)

601975	Ectodermal dysplasia/skin fragility syndrome (3)
601990 602011	Neuroblastoma (1) (?) Pancreatic endocrine tumors (1) (?)
602014	Hypomagnesemia with secondary hypocalcemia (2)
602023	Bartter syndrome, type 3 (3)
602025	Obesity/hyperinsulinism, susceptibility to (2)
602026	Refsum disease, 266500 (3)
602066	Convulsions, infantile and paroxysmal choreoathetosis (2)
602067	Cardiomyopathy, dilated, 1F (2)
602078	Fibrosis of extraocular muscles, congenital, 2 (2)
602080	Paget disease of bone-2 (2)
602081	Speech-language disorder-1 (2)
602082	Corneal dystrophy, Thiel-Behnke type (2)
602084	Endometrial carcinoma (2)
602085	Postaxial polydactyly, type A2 (2)
602086	Arrhythmogenic right ventricular dysplasia-3 (2)
602087	Arrhythmogenic right ventricular dysplasia-4 (2)
602088	Nephronophthisis, infantile (2)
602089	Hemangioma, capillary, hereditary (2)
602091	Marfan syndrome, atypical (3)
602092	Deafness, autosomal recessive 18 (2)
602094	Lipodystrophy, familial partial (2)
602096	Alzheimer disease-5 (2)
602099	Amytrophic lateral sclerosis-5 (2)
602116	Glioma (1)
602117	Prader-Willi syndrome (1) (?)
602121	Deafness, autosomal dominant nonsyndromic sensorineural, 1, 124900 (3)
602134	Tremor, familial essential, 2 (2)
602136	Adrenoleukodystrophy, neonatal, 202370 (3)
	Refsum disease, infantile, 266510 (3)
	Zellweger syndrome-1, 214100 (3)
602153	Monilethrix, 158000 (3)
602216	Peutz-Jeghers syndrome, 175200 (3)
602218	Townes-Brocks syndrome, 107480 (3)
602221	Stem-cell leukemia/lymphoma syndrome (3)
602225	Cone-rod retinal dystrophy-2, 120970 (3)
	Leber congenital amaurosis, type III (3)
602235	Epilepsy, benign, neonatal, type 1, 121200 (3)
602279	Oculopharyngeal muscular dystorphy, 164300 (3)
	Oculopharyngeal muscular dystrophy, autosomal recessive, 257950 (3)
602280	Retinitis pigmentosa-14, 600132 (3)
602363	Ellis-van Creveld-like syndrome (2)
602397	Cholestasis, benign recurrent intrahepatic, 243300 (3)
	Cholestasis, progressive familial intrahepatic-1, 211600 (3)
602404	Parkinson disease, type 3 (2)
602421	Congenital bilateral absence of vas deferens, 277180 (3)
	Cystic fibrosis, 219700 (3)
	Sweat chloride elevation without CF (3)
602447	Coronary artery disease, susceptibility to (3)

WO 01/22920 PCT/US00/26524

## 405

602460	Deafness, autosomal dominant 15, 602459 (3)
602475	Ossification of posterior longitudinal ligament of spine (2)
602476	Febrile convulsions, familial, 1 (2)
602477	Febrile convulsions, familial, 2 (2)
602491	Hyperlipidemia, familial combined, 1 (2)
602522	Bartter syndrome, infantile, with sensorineural deafness (2)
602544	Parkinson disease, juvenile, type 2, 600116 (3)
602574	Deafness, autosomal dominant 12, 601842 (3)
	Deafness, autosomal dominant 8, 601543 (3)
602575	Nail-patella syndrome with open-angle glaucoma, 137750 (3)
	Nail-patella syndrome, 161200 (3)
602616	Carbohydrate-deficient glycoprotein syndrome, type II, 212066 (3)
602629	Dystonia-6, torsion (2)
602631	Breast Cancer (3)
	Rhabdomyosarcoma, 268210 (3)
602639	Hypodontia, autosomal recessive (2)
602666	Deafness, autosomal recessive 3, 600316 (3)
602667	Nijmegen breakage syndrome, 251260 (3)
602669	Anterior segment mesenchymal dysgenesis and cataract, 107250 (3)
	Cataract, congenital (3)
602716	Nephrosis-1, congenital, Finnish type, 256300 (3)
602759	Prostate cancer, hereditary, 2, 176807 (2)
602771	Muscular dystrophy, congenital, with early spine rigidity (2)
602772	Retinitis pitmentosa-24 (2)
602782	Faisalabad histiocytosis (2)
602783	Spastic paraplegia-7 (3)

5

10

15

20

25

30

406

PCT/US00/26524

The polypeptides of the invention can be prepared in any suitable manner. Such polypeptides include isolated naturally occurring polypeptides, recombinantly produced polypeptides, synthetically produced polypeptides, or polypeptides produced by a combination of these methods. Means for preparing such polypeptides are well understood in the art.

The polypeptides may be in the form of the secreted protein, including the mature form, or may be a part of a larger protein, such as a fusion protein (see below). It is often advantageous to include an additional amino acid sequence which contains secretory or leader sequences, pro-sequences, sequences which aid in purification, such as multiple histidine residues, or an additional sequence for stability during recombinant production.

The polypeptides of the present invention are preferably provided in an isolated form, and preferably are substantially purified. A recombinantly produced version of a polypeptide, including the secreted polypeptide, can be substantially purified using techniques described herein or otherwise known in the art, such as, for example, by the one-step method described in Smith and Johnson, Gene 67:31-40 (1988). Polypeptides of the invention also can be purified from natural, synthetic or recombinant sources using techniques described herein or otherwise known in the art, such as, for example, antibodies of the invention raised against the colon cancer antigens in methods which are well known in the art.

The present invention provides a polynucleotide comprising, or alternatively consisting of, the nucleic acid sequence of SEQ ID NO:X, and/or a cDNA contained in the ATCC deposit. The present invention also provides a polypeptide comprising, or alternatively, consisting of, the polypeptide sequence of SEQ ID NO:Y and/or a polypeptide encoded by a cDNA contained in the ATCC deposit. Polynucleotides encoding a polypeptide comprising, or alternatively consisting of the polypeptide sequence of SEQ ID NO:Y and/or a polypeptide sequence encoded by a cDNA contained in the ATCC deposits are also encompassed by the invention.

Many polynucleotide sequences, such as EST sequences, are publicly available and accessible through sequence databases and may have been publicly available prior to conception of the present invention. Preferably, such related polynucleotides are specifically excluded from the scope of the present invention. To list every related sequence would unduly burden the disclosure of this application. Accordingly, for each "Contig Id" listed in

WO 01/22920 PCT/US00/26524

407

the third column of Table 7, preferably excluded are one or more polynucleotides comprising a nucleotide sequence described in the second column of Table 7 by the general formula of a-b, each of which are uniquely defined for the SEQ ID NO:X corresponding to that Contig Id in the fourth column of Table 7. Additionally, specific embodiments are directed to polynucleotide sequences excluding one, two, three, four or more of the specific polynucleotide sequences referenced by Genbank Accession No. for each Contig Id which may be included in column five of Table 7. In no way is this listing meant to encompass all of the sequences which may be excluded by the general formula, it is just a representative example. All references available through these accessions are hereby incorporated by reference in their entirety.

5

10

able 7

<b>-</b>	CONTIG	GENERAL FORMULA	ACCESSION NUMBERS
	ë		
m	390631	Preferably excluded from the present invention are one or more	AL119989, T80240, AA773747, AA809992, AA281432, AF051311, AF053535, AF070615, AF145284,
		polynucleotides comprising a	AB014560, U65313, AF145285
		is any integer between 1 to 393 of	
		SEQ ID NO:1, b is an integer of 15	
		to 407, where both a and b	
		correspond to the positions of	
		nucleotide residues shown in SEQ ID	
		NO:1, and where b is greater than	
		or equal to a + 14.	
	410299	Preferably excluded from the	
		present invention are one or more	
		polynucleotides comprising a	
		nucleotide sequence described by	
		the general formula of a-b, where a	
		is any integer between 1 to 399 of	
		SEQ ID NO:2, b is an integer of 15	
		to 413, where both a and b	
		correspond to the positions of	
		nucleotide residues shown in SEQ ID	
		NO:2, and where b is greater than	
		or equal to a + 14.	
	456200	Preferably excluded from the	N76659, T85798, AW379474, AR016730, D50857
		present invention are one or more	
		polynucleotides comprising a	
		nucleotide sequence described by	
		the general formula of a-b, where a	
		is any integer between 1 to 460 of	

			is an integer both a and b					
			correspond to the positions of nucleotide residues shown in SEO ID					
			d where b is greater than					
			- 1					
4	HCFCY21	456438	Preferably excluded from the	AI866002,	A1476046,	AI610645,	AI561299,	
			present invention are one or more	AI433976,	AI251830,	AW085667,	AW403717,	
			polynucleotides comprising a	AI868831,	AL045266,	AI801325,	AL038605,	
			nucleotide sequence described by	AW117746,	AI815855,	AI873644,	AI674838,	
			the general formula of a-b, where a	AI678599,	AI609593,	AI498579,	AI799199,	
			is any integer between 1 to 1829 of	AI867042,	AL039086,	AW051107,	AI696626,	
			SEQ ID NO:4, b is an integer of 15	AI174394,	AI521012,	AI633419,	AI890806,	
			to 1843, where both a and b	AI796743,	AW103371,	AA640779,	AW162071,	
			correspond to the positions of	AI816010,	AI480118,	AI824557,	AI499285,	-
			nucleotide residues shown in SEQ ID	AI340582,	AW059837,	AW071417,	AW132056,	
			NO:4, and where b is greater than	AI569616,	AI872711,	AI872545,	AI918655,	
			or equal to a + 14.	AI955917,	AI620003,	AI862139,	AI696612,	
				AW081036,	AI289937,	AI274508,	AI434468,	
				AI284131,	AW082040,	AW302988,	AI890833,	
				AI926790,	AI568870,	AW102785,	AW103893,	
				AI564719,	AI281772,	AI889376,	AI524671,	
				AW051258,	AI919345,	AI554245,	AI921248,	
-				AI611738,	AW002342,	AI619502,	AI677796,	
				AI632408,	AI802542,	AI308035,	AI886753,	
				AI933589,	AW026882,	AI636719,	AI476109,	
				AI923768,	AI783504,	AL079963,	AL036396,	
				AI567351,	AI620284,	AL119863,	AI500039,	
				AI274013,	AW301505,	AI922365,	AW195968,	
				AI587288,	AI345587,	AI433157,	AI702073,	
				AL036759,	AI366549,	AI446373,	AW238730,	
			•	AI500706,	AI537677,	AW083804,	AI520931,	
				AI500662,	AI633125,	AW161579,	AL041772,	
				AI648663,	AI500523,		AW005858,	
				AI284517,	AW129916,	AI242249,	AI888944,	
				AW050522,	AW192375,	AA494167,	AW268220,	

				1870781A	DT491897	AT349645	AT364788	Γ
					00001014	10101111111111111111111111111111111111		
	•			A1224392,	AI318280,	A1284509, A1.042628	ALU36146, AT673256	
				1929	ATSARORS	AI.036403	AT250663	
				5421	AA427700.	AI571909.		
	-			9131	AI922901,	AI362637,	AI924971,	
	_			AI569583,	AI554427,	AI273142,	AL040241,	
	<u> </u>			AW088134,	AI269862,	AI500553,	AI345347,	-
				AL119836,	AI612759,	AW150578,	AW190042,	
				AI269696,	AI922676,	AI800453,	AI800433,	
				AI921176,	AI888953,	AI886124,	AI499463,	
				AI874166,	A1445165,	AI963216,	AI590120,	
	-			AI308032,	AW149227,	AI828731,	AI282326,	
	-			AI590118,	AW079159,	AI287326,	AI343059,	
		•		AA572758,	AW023590,	AW169653,	AI648684,	
<del></del>				AI687065,	AI608676,	AI811845,	AI349933,	
				AI863014,	AI468872,	AI950664,	AI280661,	
				AI345608,	AL120853,	AI340603,	AI680498,	
				AI537617,	AW088903,	AI567360,	AI499381,	
				AI281779,	AI348897,	AW168650,	AI349004,	
				AW081255,	AI383919,	AI280637,	AI539153,	
				AI539771,	AI824444,	AI608936,	AI866608,	
				AI611743,	AI687362,	AI862144,	AC006373,	
				AC009501,	AC004808,	AL035407,	AC006313,	
				AC008014,	AC004470,	Z98036, A	AC004159, AC006039,	
				AL022394				
5	HMKCO08 4	467315	Preferably excluded from the	AI905893,	AI905911,	D50640, Z2	22867	
			present invention are one or more					
			polynucleotides comprising a					
-			nucleotide sequence described by					
			the general formula of a-b, where a					
			127					
			SEQ ID NO:5, b is an integer of 15					
			to 471, where both a and b					
	_		correspond to the positions of					
			nucleotide residues shown in SEQ ID		•			

			סיו	
			or equal to a + 14.	
9	HBAGS04	471563		AA148530, AA463550,
			present invention are one or more	
			polynucleotides comprising a	AC006484, Z75888, Y10196
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 891 of	
			SEQ ID NO:6, b is an integer of 15	
			to 905, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:6, and where b is greater than	
			or equal to a + 14.	
7	HALSQ75	488131	Preferably excluded from the	AI352096, AA376070, T81033
			present invention are one or more	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 398 of	
			SEQ ID NO:7, b is an integer of 15	
			to 412, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:7, and where b is greater than	
8	HMVBD21	490848	Preferably excluded from the	AI767324, AL121194, AA972628, AI095851,
			present invention are one or more	AA743343, AW366882, D20570, AC009802
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 738 of	
			SEQ ID NO:8, b is an integer of 15	
			to 752, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	

HKIMD67 HOOAE34 HHSDD62	500696	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 628 of SEQ ID NO:9, b is an integer of 15 to 642, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:9, and where b is greater than or equal to a + 14.  Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 197 of SEQ ID NO:10, b is an integer of 15 to 211, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:10, and where b is greater than or equal to a + 14.  Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a	T71949, N95702, AI306688, AW451579, AI341434, AI223407, AA885055, AA846712, AA379446, AW362461, H42165, D80959, AI928895, AA081721, D87459, AF134303 AI380563, AI004009, AI625234, AW337321, AA321125, AA937785, AA363438 AM409739, AA773074, AI083705, AI870827, AI126674, W94001, AI273489, AI275482, AI264045, AI126674, W94001, AI273489, AI275482, AI350937, AI350937, AI393684, R76773, AA483544, AW131780, AI091146, AA130101, AA255799, AA976695, AA774090, R76460, AI206751, AI350938,
		is any integer between 1 to 518 of SEQ ID NO:11, b is an integer of 15 to 532, where both a and b correspond to the positions of	, AI669701, AA669454, H04098, , R49064, AA130100, AA639898, , AA082299, AA702716, X07233,

			NO:11, and where b is greater than	
12	HSLGZ32	506619		AA316122, AA314900, AF121202
			present invention are one or more	
			polynucleotides comprising a	
			the general formula of a-b, where a	
			is any integer between 1 to 1106 of	
			SEQ ID NO:12, b is an integer of 15	
			to 1120, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:12, and where b is greater than	
			or equal to a + 14.	
13	HCENL15	507852	Preferably excluded from the	N40063, AA233205, R46529, AI015135, AW130559,
			present invention are one or more	AA324511, AI460380, AA346401, AA604942,
			polynucleotides comprising a	AI125644, AI703464, T67213, AW103052, AI452537,
			nucleotide sequence described by	AI050784, AI949725, AI052071
			the general formula of a-b, where a	
			is any integer between 1 to 586 of	
			SEQ ID NO:13, b is an integer of 15	
			to 600, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:13, and where b is greater than	
			or equal to a + 14.	
14	HCQAI38	509423	Preferably excluded from the	Ì
			present invention are one or more	AW274753, T35439, T10802
			polynucleotides comprising a	
			nucleotide sequence described by	-
			the general formula of a-b, where a	
			is any integer between 1 to 793 of	
			SEQ ID NO:14, b is an integer of 15	
			to 807, where both a and b	
			correspond to the positions of	
		-	nucleotide residues shown in SEQ ID	

			NO:14, and where b is greater than or equal to a + 14.	
15	HPMDT48	509734	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 402 of SEQ ID NO:15, b is an integer of 15 to 416, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:15, and where b is greater than or equal to a + 14.	
91	HADFX66	509856	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 738 of SEQ ID NO:16, b is an integer of 15 to 752, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:16, and where b is greater than or equal to a + 14.	AA845353, N31960, AA617724, AA307653, AA862795, N92883, R60191, AA321715, AI802963, W23451, AA905145, W25563, AA469079, R60190, R18173
17	HONA101	524721	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 467 of SEQ ID NO:17, b is an integer of 15 to 481, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID	AA362921, AA436291, AA425301, AL079672, AW361081, AW341687, AA284486, AI880015, AI379662, AI935013, AA436164, AI066555, AA837415, AA706542, AI126021, AB028996

			NO:17, and where b is greater than		
			or equal to a + 14.		
18	HEBBT54	524901	Preferably excluded from the	T85603, AA132177, AA314644, AA1	AA133670, T99921,
			present invention are one or more	AL039938, R01637, AA046158, APC	AP000497, D88153
			polynucleotides comprising a		
			nucleotide sequence described by		
			the general formula of a-b, where a		
			is any integer between 1 to 898 of		
			SEQ ID NO:18, b is an integer of 15		
			to 912, where both a and b		
			correspond to the positions of		
			nucleotide residues shown in SEQ ID		u Paraba di
			NO:18, and where b is greater than		
			al to a + 14.		
61	H2CBG63	527600	Preferably excluded from the	AA307234, R60594, AC007327	
			present invention are one or more		
			polynucleotides comprising a		
			nucleotide sequence described by		
			the general formula of a-b, where a		
			is any integer between 1 to 493 of		
			SEQ ID NO:19, b is an integer of 15		
			to 507, where both a and b		
			correspond to the positions of		
			nucleotide residues shown in SEQ ID		
			NO:19, and where b is greater than		
			or equal to a + 14.		
20	HHSBA79	527827	Preferably excluded from the	AI915163, AI690026,	AC004817,
			present invention are one or more	AL132774, AL078602, AC003999, AL136168,	AL136168,
			polynucleotides comprising a	AL022396, 'Z68325, AL035414, AB020868,	.020868, U95740,
			nucleotide sequence described by	AC007685, AL049712, AP000067, AC002992,	AC002992,
			the general formula of a-b, where a	AL022148, U95741, U95743, AC007304, AC002390,	7304, AC002390,
			is any integer between 1 to 396 of	AC005482, Z82188, AC005477, AC005295, AL035408,	:005295, AL035408,
			SEQ ID NO:20, b is an integer of 15	AC007028, AL035608, AC004917, AL132992,	AL132992,
			to 410, where both a and b	AC002385, AC007064, AC004825	
			correspond to the positions of		
			nucleotide residues shown in SEQ ID		

			NO:23, and where b is greater than or equal to a + 14.	
24	HSAAX52	530773	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 447 of SEQ ID NO:24, b is an integer of 15 to 461, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:24, and where b is greater than or equal to a + 14.	AA299283, AW379368, AA374069, D61135, D60581
25	НАССЕ33	532810	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 439 of SEQ ID NO:25, b is an integer of 15 to 453, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:25, and where b is greater than or equal to a + 14.	AA305030, AI207985, AA469325, AA420424, D10040, L09229
26	HE8DA85	533242	preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 1926 of SEQ ID NO:26, b is an integer of 15 to 1940, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID	AI052713, AA993209, AI133542, AI160185, AI110772, AA443423, AA406485, AI806057, AI1114495, AA410346, AI436310, AI370818, AL035763, T11697, C21213, AA707071, N54577, H52269, R89600, F06943, R98692, R97937, W84760, R98918, AA358620, AI207561, H65229, AI123345, H60435, R98546, AA010003, R98713, AA332857, AI131251, T75531, H52507, T29077, AW440733, R98714, T53093, H80150, H78893, H79897, AA010004, R20093, W84755, AI351429, AI185625,

			NO:26, and where b is greater than or equal to a + 14.	A628869, N76658, T754 A026133, AA342031, H6 79803, N77400, T81302 L035762, T11698, T523 I056662, T52412, AA69 56352, AF086786, M152 83821, AL020991, AF06
27	HSKII86	541126	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 850 of SEQ ID NO:27, b is an integer of 15 to 864, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:27, and where b is greater than or equal to a + 14.	AA305909, AA354725, AA296543, AA347633, AA100063, AA424070, AA296485, AI795779, AW239153, AA337079, T87056, AA263171, AA044017, AA044192, AI683358, AW363341, AW138402, AA294979, AA424397, AW134673, T75498, AA827350, M55542, M55543, AR035947, M55544, M63961, AF109168, AJ007970, AF077007, M80367, AR035948
28	HUSG139	542268	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 689 of SEQ ID NO:28, b is an integer of 15 to 703, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:28, and where b is greater than or equal to a + 14.	AW407143, AA213542, AA284733, AA485799, AI761438, AC005368
29	HKIMB44	547920	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a	A1985978, AW070887, R76142, AW089184, N31775, AI687598, AI963830, AL035869, AI697270, R83410, W94030, AW083686, AI188661, AA653485, AA165514, AW188780, AA989157, AI473469, H69762, AI925548, AA908930, AA340369, AI682723, AI950093,

		is any integer between 1 to 323 of	AW438743 AT654303 H70528 T82154 H63221
		NO:29, b is an integer of	AI186315, W85840, W26638, AW190
		both	, AA961655, D55762, D55763, D54646,
		correspond to the positions of	AA089984,
		nucleotide residues shown in SEQ ID	C15803, AA1655
		NO:29, and where b is greater than	9, AW194507, AI110844
		oa + 14.	AA100718, AI821986, AI085242, AA903287,
			, AI536908, AA053088,
			AI821788, AI821745, T52103, AI540625, AA011625,
			AA572953,
			AA284416, J05032, AC003666, Z69838, AC005952,
			AL031679, AL024474, AC000112, AP000361,
			AL021977, AC005215, AC004903, AC006238,
			AF113009, AF095901, AL049636, AC004551, Z99289,
			0, AC003104
			AP000214, Z81310, AP000255, AC004851, AP000135,
			, AC007245
			, AC005520, AC005154,
			_
			AL034418, AC002382, AC005037, AC007240,
			4, AC005018, AF207550,
			~
			AC005999, AL132641, Z84488, AF001549, AC006453,
			AL008639, AC004099, AC007463, AC008163, Z98044,
			AC007387, AF141976, AC002365, L30117, U73023,
			AL033543, AC007160, AC003119, AL078581
30 HBMVJ62	549642	Preferably excluded from the	AI523366, AA856766, AI690026, AA856757,
		present invention are one or more	AA486994, AA491240, AA214142, AI032325, R10942,
		polynucleotides comprising a	AW023137, R10519, AW237618, AL049712, AC007388,
		nucleotide sequence described by	AC007002, Z68192, AC004104, AC007425, AC007182,
		a-b, when	2, AL132774
		r betw	
		NO:30, b is an int	, AL121840,
		ere both a and b	AC005291, AF002994, AC006991, AC002394,
		correspond to the positions of	AB020863, Z82210, AP001058, AC005609, AC004388,

			nucleotide residues shown in SEQ ID NO:30, and where b is greater than or equal to a + 14.	AC001526, AL031073, AL049828, U69730, AC005057, Z69648, AC007685, AL096861, AL079304, AC005011, AL022578, AC004858, AC008064, AF109907, AP000338, AP000216, AL136168, AC005821, AC003964, AB020868, AC007656, AC007437, AC006210, AL035414, AC004103, AC005971, AP000067, AC005697, AC009946, AC005971, AC002331, Z97054, Z82188, AC004791, AC005380, AC005616, AC004679, Z97055, AC003991, AC007263, AC007280
31	HBXFC78	550207	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 557 of SEQ ID NO:31, b is an integer of 15 to 571, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:31, and where b is greater than or equal to a + 14.	1, A1689429 3, N70051, A1291826, 5, N50944, 0, A1783813
32	HE2FR32	552115	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 410 of SEQ ID NO:32, b is an integer of 15 to 424, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:32, and where b is greater than or equal to a + 14.	C19093, AA479586, AI417611, AA477425, W07367, AW403461, AA292357, N32437, AA258489, U69127
33	HKACD58	552465	LS 51	AA877796, AW027434, AI335269, N46240, AW402301, AI525602, H38504, AW390227, AI365603, AI819188,

 polynucleotides comprising a	AW390207, AW368379, R53778, AA370005, AA134615,
sedne	AA354527, R69656, AI524965, AA425001, F37313,
$\vdash$	374
eger between 1 to 1612	R78049, AA355898, AI360437, AA149032, R55146,
SEQ ID NO:33, b is an integer of 15	04915, R881
to 1626, where both a and b	, AI193372, AA380842,
 correspond to the positions of	8997, AI096656, AI367032,
	5242, AA149757, AI433008
NO:33, and where b is greater than	AA555145, T27702, AI249880, AI434080, AI561147,
 or equal to a + 14.	i, AL046021,
	AI732975, AL110373, AA853473, AI805349,
	AA903287, R46841, AA551390, AA662117, AI821062,
	AI368745
	AW081103, AA019257, F00107, AL046262, AI557808,
	AI064787, AA618452, AI110708, AL040077,
	AA653459, AA764903, AI627181, AA235975,
	AI823716, AA877935, AA659232, AI821259,
	AI887241, AL110402, AW089171, AA937752,
	AI874222, AL138455, N52358, AW079659, AI133029,
	AI111171, AA806438, AI345797, AL042753,
	AI
	7,
	AI821788, AI821745, AI207728, AW104715,
	9447, AI039141, AI369580,
	6, AW160760, AL038134,
	2, AL050037, AC006455,
	, AL031282,
	211, AP000133, AP000030,
	520, AC006017, AF095901,
	, AC004990, AF003738, V
	AP000361, AL136130, AC002078, Z84489, AC007748,
	C004974, AC0050
	C006458, AL031274, Z82206,
	44, AC009044, AC005284
	C006205, J00349, Z98036, AL031774, A
	AC011013, AC005006, AC005078, AC002382,

	42
	AC005074, AF090940, L30117, J05032, AF045527,
	, AF057280, AF126531, X91656,
	, Z82250, AC
	AC004209, AC000394, AC003005, AC006139,
	Α,
	AL031662, AF024533, AC005353, Z99714, Z93784,
	AC002564, AL022315, AC002467, AP000206,
	AC002301, AL049761, Z98748, AL031656, AC004900,
	ò
	_
	AC002432,
	_
	4, AC004544
	60, Z99289, AC005296,
	AP000319,
	7, AC000053,
	AP000569, AC005057, AC007216, AL050309,
	AL031346,
	, AC002482, AL079340,
	86, AC002086
	AC004589, Z49235, AF195658, AC004210, AC002559,
	1, AP000508, AC0024
	553, Z92543, Z8
	4, AC006029, AC007671,
	086, AC005209
	, D843
-	465, Z99297, AB012260, U7
	-
	AC005992, D83989, S75201, AF088219, AC006582,
	989, AB012179, AC004972,
	3, AL079342,
	., AC000025, X581
	87, Y11740, Z49258, AC004760, AC00
	AL035450, AC004940, AL080245, X66401, AC004769,

				AC006221, AC004009, AP000338, AL022399, AL022393, AL033523, AC005008, AC005250, AL096770
34 4	HDAAB62	5 5 4 3 6 9 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 436 of SEQ ID NO:34, b is an integer of 15 to 450, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:34, and where b is greater than or equal to a + 14.	AA307995, AA312596, AA307928 AA307129, AA102186, AA343955 AA361582, AA362045, N86980, AL046225, AI241561, N66944, 227098, AL045756, C17672, AA A1673070, AA622421, AA662921 AA769512, AA587021, AA151746 N68288, M81691, AI821039, AI36 AA196412, X16396, AL024507, AC00378, AL135783, AC005328 AL031666, Z82198, D13631, D2 AC00378, AL135783, AC005328, AC003072, Z98256, AL031320, AC005229, AL109758, U40369, AC005239, X55448, AC002352, AC005398, X55448, AC002352, AC005899, AL035398, Z84469, AC005899, AL078602, AC003102, AC005899, AL078602, AC003102, AC002425
8	HEPBA24	557152	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 946 of SEQ ID NO:35, b is an integer of 15 to 960, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:35, and where b is greater than or equal to a + 14.	AA632191, AW151795, R59316, AI338706, AI276888, AI366798, AI471791, AA565616, AI248949, AI269722, AI393882, AA504204, AA813608, AI335657, AA682615, AI039562, AA400139, AA187165, AI376431, AA454074, AA128383, AI022866, AA929034, R46374, AA810231, AI241427, AA865170, AI567959, W93122, N47805, AA35391, AI919230, AI242499, AA401552, T28382, AA453654, AA33614, AA336149, AA335946, R41308, T24710, AI017254, D26018

36	HOGBL08	557230	Preferably excluded from the	AA151092, AA367221, F09233, AA442256, AA442255,
			present invention are one or more	AA933632, W80475, H22100, AI274366, AA909429,
			polynucleotides comprising a	AI918108, AI499770, AI278711, AA948413,
			nucleotide sequence described by	AA831784, AI470908, AA936238, W94676, AI002684,
			the general formula of a-b, where a	AI050788, AI264677, AI246792, AI276574,
			is any integer between 1 to 516 of	AI352684, AA428287, AI952002, AI890744,
			SEQ ID NO:36, b is an integer of 15	AI806291, T18857, AI819539, R43330, AI749667,
			to 530, where both a and b	R71480, AI262455, N75456, AI869496, AA826541,
			correspond to the positions of	AA436899, AI187706, AL135960, AJ131016, U21049,
				AC006487, U80460
			NO:36, and where b is greater than	
			or equal to a + 14.	
37	HCYBD62	558366	Preferably excluded from the	AA305096, AA424823, AA995932, AW001055,
			present invention are one or more	AI016665, AI399850, AI636182, AI949894,
			polynucleotides comprising a	AW361640, AI634640, AW377132, AW377117,
-			nucleotide sequence described by	AW388099, W07829, AA993439, X89602, X67098
			the general formula of a-b, where a	
			is any integer between 1 to 524 of	
			SEQ ID NO:37, b is an integer of 15	
			to 538, where both a and b	
			correspond to the positions of	
			NO:37, and where b is greater than	
			o a + 14.	
38	H2CBD20	961015	Preferably excluded from the	AA307235, AI002535, H49502, AL110292
			present invention are one or more	
			بيد	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			eger between 1 to 1242	
			SEQ ID NO:38, b is an integer of 15	
			to 1256, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:38, and where b is greater than	
			or equal to a + 14.	

HETDN09	573181	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 652 of SEQ ID NO:39, b is an integer of 15 to 666, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:39, and where b is greater than or equal to a + 14.  Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 1002 of SEQ ID NO:40, b is an integer of 15 to 1016, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:40, and where b is greater than or equal to a + 14.	AI761465, AW270500, AA290850, AA953717, H57392, AI127126, T82331, AW293000, AA973625, AI298354, AW451105, Z64718, Z64717  AI022684, AI807626, AI032750, AI026655, AA931238, AI806336, AI68363, AA062961, AA931238, AI806336, AI68363, AA062961, AA035619, AA701296, AW351549, AA613277, AA918186, AI500592, AA812859, AI339340, AA911056, AI13829, AA962373, AI826452, AI620160, AI13829, AA962373, AI826452, AI620160, AI31126, AA962571, AI20552, AI139296, AI311267, AI310838, R71902, AI311267, AI310838, R71902, AI311267, AI34075, AI34054, AA803452, AC008132, AV311175, AI34075, AI340244, AB000356, AC008132, AV000550, AC008018, AC011718, AC007981, AC002308, M30474, AP000354, XI5443, M33822, M33821, AC007325, AI0399, L10399, L10396, Z93346, AC00334, AC007380, Z93344, Z93344, Z93343, AU07380, Z93344, Z93343, AU07380, Z93344, Z93343, AU07380, Z93344, Z93343, AU07380, Z93344, Z93347, AU07380, AU07380, Z93344, Z93347, AU07380, Z93342, Z93344, Z93347, AU07380, Z93342, Z93344, Z93347, AU07380, Z93342, Z93344, Z93347, AU07380, Z93342, Z93344, Z93347, Z93348, Z93347, Z93348, Z93427, Z93448, Z93448, Z93448, Z9
HCYBE04	573793	Preferably excluded from the	AJ006789 AA305129, AA768244, AA310241
		are one or	

			otides comp	
			al formula of a-b, wher	
			is any integer between 1 to 409 of	
			SEQ ID NO:41, b is an integer of 15	
			to 423, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:41, and where b is greater than	
			or equal to a + 14.	
42	HDPFI14	573796	Preferably excluded from the	AA316491, AA704220, R80096, AA305136, R57983,
			present invention are one or more	ω ω
			polynucleotides comprising a	D89852, D89851
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 947 of	
			SEQ ID NO:42, b is an integer of 15	
			to 961, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:42, and where b is greater than	
			or equal to a + 14.	
43	HJBCD90	574094	Preferably excluded from the	AA311657, N88312, AA307717, Z21301, AA551523,
			present invention are one or more	4
			polynucleotides comprising a	D80391, D59787, D80439, D80196, AA305409,
			nucleotide sequence described by	3, D51799, D59859, D51060, C15076,
			the general formula of a-b, where a	D80166, C14014, D80195,
			is any integer between 1 to 531 of	2, D80210, D80240,
			SEQ ID NO:43, b is an integer of 15	D50995, C14331, D59467,
			to 545, where both a and b	D80038, D80043,
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	D80366, D80188, D80248, AA514186, D50979,
			NO:43, and where b is greater than	D51103, AA305578, D80157, D80241, D80193,
			or equal to a + 14.	D80045, D59889, D5
				, D81111, C03092, AW377671, D8
				D59551, AW177440, AW178893, C14429, AI525923,

	D80251. F13647. D59503. AA809122. C14227.
	11417, D58101, AW375405, AW360844, TO
	59317, H67854, H67866, AW360811, C14973,
	344, AW360817, AW360834, D582
	D80258, D80064, D
	l, AI535686, <i>I</i>
	AW179328, D59695,
	76, AW375406, AW378534, AI525
	AW352171, AW179332, AW377672, AW179023,
	178905, AW179013, Z21582, AW17773
	2,
-	019, AW360841, AW179024,
	20, AA51418
	C14407, AW378539,
	AW178909, H67858, AW17745
	, AW352170, AW178986, AI5252
	3, AW178908, AW179
	D31458, AW17
	5912, AA285331, AW177734, A
	9017, C06084, C13958, AW367950, C1
	80168, AW178759, AW179009, AW179
	52, T02974, AI525237, AI535665
	3542, C05763, D52291, C14298, D512
	78911, C14077, AW378525, AW352163, AW
	868, AW177728, T03048, AW369651, AI52522
	91, AI525914, AC007262, A8
	), AJ132110, AR018138, F
	00, A62298, AR008278, AF058696,
	AR054175
	, AR0601
	, A94995, D26022, AR008277,
	5, AR052274, A43192, Y12724, A6
	0, AR038669, A25909, AR066488,
	066487, A70867, A67220, D89785, A7
	D34614, A30438, AR062872, AR008443, AR016691,

				DR016690 179511 1146128 B64136 B68321
				88547, D50010, X68127, X8 AR025207, AF123263, AR06
44	HAJAB40	574927	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 363 of SEQ ID NO:44, b is an integer of 15 to 377, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:44, and where b is greater than or equal to a + 14.	AA316028
45	H2MCA74	575139	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 426 of SEQ ID NO:45, b is an integer of 15 to 440, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:45, and where b is greater than or equal to a + 14.	AA112812, H73681, AW L035458, AC Z21853, AC AC006369, AL122127, AF017732, AP000346, AL021155, AF017188,
46	HWBAX42	575591	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 511 of SEQ ID NO:46, b is an integer of 15 to 525, where both a and b correspond to the positions of	AW068735, AA853585, AA380263, AL121408, AL121410, X64330, U18197, L27075, U05210, L47278

			-	نداحات المراجعة المرا
			nucleotiae restaues snown in aby in	
			<b>⊢</b> i	
			or equal to a + 14.	The second secon
47	HLMMR55	576132	Preferably excluded from the	
			present invention are one or more	
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 400 of	
			SEQ ID NO:47, b is an integer of 15	
			to 414, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:47, and where b is greater than	
			or equal to a + 14.	
48	HNFGN91	577390	Preferably excluded from the	AJ011930
			present invention are one or more	
			polynucleotides comprising a	
-			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 309 of	
			SEQ ID NO:48, b is an integer of 15	
			to 323, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:48, and where b is greater than	
			or equal to a + 14.	
49	06IQMJH	577685	Preferably excluded from the	AA769584,
			present invention are one or more	AA970935, AI807551,
			polynucleotides comprising a	AA769047, AA804530, AW340028, AA286746
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 827 of	
			SEQ ID NO:49, b is an integer of 15	
			to 841, where both a and b	
			correspond to the positions of	

			- 1	
			NO:49, and where b is greater than or equal to a + 14.	
	HCQAB18	578079		AA721676, AI632745, AI478171, AI719338, AA761073, AI766631, AA485859, AW059674, AI797505, AI709367, AC000123, AC000127
51	HELHI45	578660	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 303 of SEQ ID NO:51, b is an integer of 15 to 317, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:51, and where b is greater than or equal to a + 14.	N36929, AA771779, AW196937, AA342301, AI808034, AI432219, AI694329
52	HNHDV16	280860	i i i i i i i i i i i i i i i i i i i	AL049874, Z84488, AC002549, AC003035

			uncleotide residues shown in SEO ID	
			NO:52, and where b is greater than	
			or equal to a + 14.	
53	HOAAD32	581143	Preferably excluded from the	AA307601, AI760475, AI972520, AI990288,
			present invention are one or more	AI675118, AI669210, AW003506, AW016052,
			polynucleotides comprising a	AI880850, AA807606,
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 640 of	, AI300737,
			SEQ ID NO:53, b is an integer of 15	5, AI830638
			to 654, where both a and b	AI342005,
			correspond to the positions of	R33457, AA935544, AI738445, AA648791, AI625335,
			nucleotide residues shown in SEQ ID	R68590
			NO:53, and where b is greater than	
			or equal to a + 14.	
54	HSAVM80	584899	Preferably excluded from the	AI902580
			present invention are one or more	
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 320 of	
			SEQ ID NO:54, b is an integer of 15	
			to 334, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:54, and where b is greater than	
			or equal to a + 14.	The state of the s
55	HWLMA5	699009	Preferably excluded from the	AI341167, AI652526, AI990232, Z22968, Z22969,
			present invention are one or more	Z22971, Z22970, Y18391, Y18392, Y18390
		_	polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 460 of	
			SEQ ID NO:55, b is an integer of 15	
			to 474, where both a and b	
			correspond to the positions of	

			nucleotide residues shown in SEQ ID			The state of the s
			•-			
			or equal to a + 14.			- 1
56	HE8BQ01	611839	Preferably excluded from the	AI267198,	AW242820,	ο
			present invention are one or more	AA034283,	AI912188,	AA2825
			polynucleotides comprising a	AI955322,	AI680802,	-
			nucleotide sequence described by	AW340279,	AI633670,	AI140173, AI914144
			the general formula of a-b, where a			
			is any integer between 1 to 353 of			
			SEQ ID NO:56, b is an integer of 15			
			to 367, where both a and b			
			correspond to the positions of			
			nucleotide residues shown in SEQ ID			
			NO:56, and where b is greater than			
			or equal to a + 14.			
57	HELHD03	614078	$\neg$	AW028557,	AW008015,	AA279640
			present invention are one or more			
			polynucleotides comprising a			
			nucleotide sequence described by			
			the general formula of a-b, where a			
			is any integer between 1 to 550 of			
			SEQ ID NO:57, b is an integer of 15			
			to 564, where both a and b			
			correspond to the positions of			
			nucleotide residues shown in SEQ ID			
			NO:57, and where b is greater than			
			or equal to a + 14.			
58	HBMCT70	614554	Preferably excluded from the	N31002, A	AA504707, AJ	AL041182, M78574
			present invention are one or more			
			polynucleotides comprising a			
			nucleotide sequence described by			
			the general formula of a-b, where a			
			is any integer between 1 to 430 of			
			SEQ ID NO:58, b is an integer of 15			
			correspond to the positions of			

			nucleotide residues shown in SEQ ID				
			or equal to a + 14.				
59	HLYDF04	615029	Preferably excluded from the	AI972404,	AW172842,	AL079983,	AI769801,
			present invention are one or more	AI769431,	AI472252,		D79445, AI282586,
			polynucleotides comprising a	AI935375,	AI569290,	AA450323	
			nucleotide sequence described by				
			the general formula of a-b, where a				
			is any integer between 1 to 333 of				
			SEQ ID NO:59, b is an integer of 15				
			to 347, where both a and b				
			correspond to the positions of				
			nucleotide residues shown in SEQ ID				
			NO:59, and where b is greater than				
			or equal to a + 14.				
09	HDSAP04	615590		AI732729,	AI858825,	AI625874,	AI266164,
			present invention are one or more	AA402921,	AA426648,	AI039796,	AI567237,
			polynucleotides comprising a	AW090009,	AC000064,	AC007566	
			nucleotide sequence described by				
			the general formula of a-b, where a				
			is any integer between 1 to 308 of				
			SEQ ID NO:60, b is an integer of 15				
			to 322, where both a and b				
		-	correspond to the positions of				
			nucleotide residues shown in SEQ ID				
			NO:60, and where b is greater than				
			or equal to a + 14.				
19	HWBFZ21	630230	Preferably excluded from the	AW369648,	AI904452,	AF098799,	AL137335
			present invention are one or more				
			polynucleotides comprising a				
			nucleotide sequence described by				
			the general formula of a-b, where a				
			is any integer between 1 to 820 of				
		_	SEQ ID NO:61, b is an integer of 15				
			to 834, where both a and b				
			correspond to the positions of				

			nucleotide residues shown in SEQ ID NO:61, and where b is greater than or equal to a + 14.	
62	<b>НС</b> QВН72	637548	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 1782 of SEQ ID NO:62, b is an integer of 15 to 1796, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:62, and where b is greater than or equal to a + 14.	AA640538, AA649644, AA649707, R31618, AA652004,
63	негсн31	637605	preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 1362 of SEQ ID NO:63, b is an integer of 15 to 1376, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:63, and where b is greater than or equal to a + 14.	1967961, AA888070, AA653693, A AI201196, N66161, AW117713, A 34009, H98804, AA969760, AI017 AW087505, R77266, R67143, AA3 N66135, AA300547, C16517, H01 D12299, H99892, R77313, N5712 25024, AA633387, N98686, AA334 12297, AI217112, AA724744, AA4 R66409, C16522, C16404, H8799 A280332, AL035410
49	HNHEU34	638125	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 560 of SEQ ID NO:64, b is an integer of 15 to 574, where both a and b correspond to the positions of	AC004876, Y12661, M60522, M60525, M74223

			nucleotide residues shown in SEQ ID		
			NO:64, and where b is greater than or equal to a + 14.		
65	HJMAF30	638188	Preferably excluded from the	AI090108, AA810218	, AA513307, AW028090,
			present invention are one or more	AI572270, AW451013	, AI373062, AI492435,
			polynucleotides comprising a	AI141965, AI161216	i, AI952357, AI420596,
			nucleotide sequence described by	AI804945, AI123032	_
			. formula	AI984276, AA588635	, C15098, AI
			is any integer between 1 to 589 of	AA902480, AI611205	, AI928306,
			SEQ ID NO:65, b is an integer of 15	AA368086, AA731886	6, AA358722, AA732765,
			to 603, where both a and b	AW383478, AA470116,	i, AI928316, AW383528,
			correspond to the positions of	AA358723, AW383489,	), AL046849, AI075391,
			reg	AA974899, AI630005,	i, AC008126
			NO:65, and where b is greater than		
			oa + 14.		
99	HWBBK93	638249	Preferably excluded from the	AI188389, AI763238	3, AI188787, AA479523,
			present invention are one or more	AI423154, AI346520	, AW005918,
			polynucleotides comprising a	AA781395, AI394520,	), AI700522, AA778127,
			nucleotide sequence described by	AW338995, AI348157,	AI139369,
			l formula	AW291338, AI061441	, AI928073,
			er between	เก	, AA430584
			SEQ ID NO:66, b is an integer of 15	AA576421, T34153,	AA418766, AI272728, AA563878,
			to 1772, where both a and b	AA528124, AI050707,	7, AA433929, AI218374,
			correspond to the positions of	AA150587, AI422109,	), AI075212, AI338694,
				AA418722, AA969029,	), AA505360, AA782389,
			NO:66, and where b is greater than	AI348193, AA662690	), AA904379, AI743240,
			oa + 14.	AI126395, AI090901	
				AA465703, AA505543,	AA771799,
				AW271730, AI248295,	AA215970,
				AA971867, AI333671	l, AI380868, AA574396, R35114,
				N33899, AW194812,	AI280563, R16040, D45470,
				AW195236, R24139,	R79672, W30758, R18084,
				_	AI365169, AI864209, N98221,
				Z40957, T85033, AA	AA618381
				AI926895, R02608,	R42978, N79943, Z45240,
				AA320674, N71980,	AI630495, T30798, AI886691,

				AW089148, R49502, R01668, R24146, AA650492, R79861, AI589880, AA705913, AI247102, AI569369,
29	HFXAK32	638319	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 1815 of	18, AI751557, R55888 , H65732, T79985, AB , AL021154, AL034429 50, AC006948, AC0065 82, AF222686, AP0006
			SEQ ID NO:67, b 1s an integer of 15 to 1829, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:67, and where b is greater than or equal to a + 14.	
89	HUSIT18	651380	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 1674 of SEQ ID NO:68, b is an integer of 15 to 1688, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:68, and where b is greater than or equal to a + 14.	AW294097, AI279800, AA316672, N25621, N34219, AA115172, AI125602, W16706, AA228116, AI348328, AA322714, R59092, AI699089, R51246, AI086372, AA383008, AW339394, AA344347, AA227730, R51245, AB011123
69	HMWBH5	651876	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 551 of SEQ ID NO:69, b is an integer of 15 to 565, where both a and b correspond to the positions of	AI554920, AI469981, AI539465, AA425263, T28976, AA812162, AI740449, W21091, AW242010, W23635, AW275702, AA476848, AI808924, AW117295, AI148939, AA811229, AI343010, AI630793, I96178, A50906, L00049, X85754, U09793, K01912, Z12125, E00392, E00393, I01060, M35504, M35505, X02456, X00485, L00047, X02454, U76425, U76426

			NO:69, and where b is greater than				
			or equal to a + 14.				
70	HCQAW11	653175	Preferably excluded from the	AL041795,	AI590066,		AA126363,
			present invention are one or more	AA463388,	AA767754,	AA463880,	AA886811,
			polynucleotides comprising a	AW079539,	AI884597,	H23284, AA	H23284, AA907711, AA789127,
			nucleotide sequence described by	AI636922,	AI039001,	AI681335, AA126259	AA126259,
			the general formula of a-b, where a	AL050120,	AC004925		
			is any integer between 1 to 661 of				
			SEQ ID NO:70, b is an integer of 15				
			to 675, where both a and b				
			correspond to the positions of				
			nucleotide residues shown in SEQ ID				
			NO:70, and where b is greater than				
			or equal to a + 14.				
71	HPRAS01	655544	Preferably excluded from the	AA370716,	AA888839,	AI817753,	AW131538,
			present invention are one or more	AA860117,	AI479976,	AI568675,	AI922252,
			polynucleotides comprising a	AW084473,	AA370730,	AA740955,	AA935921,
			nucleotide sequence described by	AI566265,	AI092718,	AA854646,	AA724492,
			the general formula of a-b, where a	AW439983,	AL109984		
			is any integer between 1 to 256 of				
			SEQ ID NO:71, b is an integer of 15				
			to 270, where both a and b				
			correspond to the positions of				
			nucleotide residues shown in SEQ ID				
			NO:71, and where b is greater than				
			or equal to a + 14.				
72	HWBBC13	656722	Preferably excluded from the	Z69042			
			present invention are one or more				
			polynucleotides comprising a				
			nucleotide sequence described by				
			the general formula of a-b, where a				
			is any integer between 1 to 524 of				
			SEQ ID NO:72, b is an integer of 15				
			to 538, where both a and b				
			correspond to the positions of				

		nucleotide residues shown in SEQ ID NO:72, and where b is greater than or equal to a + 14.	
HNTBM67	659801	excludention ides caper formulates caper formulates being and being to the tresidum where	AL134955, AA307472, M78461, D56412, AW382561 AW382555, AW382532, AW382562, AW382559, AA223890, AI536122, AA826373, AW382553, AB014509, AB011159, D84346, X80029
HDPKC15	660020	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 626 of SEQ ID NO:74, b is an integer of 15 to 640, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:74, and where b is greater than or equal to a + 14.	AL037983, AL046549, AA630672, AL137998, AA829036, T92347, AW022608, AA570230, AL041706, AW294985, AA629872, T41259, AA588001, AA700032, AA746659, AI002744, AA346467, AI907530, H25921, T41354, AA812141, AL120086, AA570797, AA601125, AA568198, AL047429, AL138182, AI249473, AA515462, AA515443, AI619436, R99735, AI922803, T47572, AC005034, AL133163, AJ006997, AL133244, AL009174, AC003036, AL034386, AC007193, AC004217, AL080317, AL034386, AC002067, AP000690, Z84487, AL121603, AC006120, AC007919, AC02485, AC005031, AC009498, AC004540, AL021367, U95740, U80017, AP000251, AL031848, AF029308, AC006042, Z69917, AC004856, AL121748, AL00984, AL034350, Z82194, AL023284, AC005207, AC00052, AC004019, U66083, AF091512, AC006262,

				AP001051, AC004967, AC005332, AC005538,
				AL109854, AP000694, AC005940, AC007011, Z81450,
				AC004652, AC005202, AL049544, AF196970, Z84476,
			-	, AFIZO4U3
75	HMAHP16	009199	excluded from the	, AL135338, A1682468,
				AA911190, AA923049,
			rides comp	~
			nucleotide sequence described by	6, N28500, A1458578, AI168801, AWO
				AI276029, AI753280, T63417, R99573, N52383,
			is any integer between 1 to 493 of	AA350643
			SEQ ID NO:75, b is an integer of 15	
			to 507, where both a and b	
			correspond to the positions of	
			res	
			NO:75, and where b is greater than	
			oa + 14.	
9/	HCE1D45	664481	Preferably excluded from the	AI633731, AA742535, AA928586, AW024580,
			present invention are one or more	AI031748, AA287493, AW139368, AA235073,
			polynucleotides comprising a	AI342861, AA024783, AI122951, N47975, AA235180,
			nucleotide sequence described by	AA936068, AW083549, AA953192, T96536, AW297014,
			the general formula of a-b, where a	), R68510, AA287388
				R75691, AI206677, C05105, AW372944, Z41074,
			SEQ ID NO:76, b is an integer of 15	N89741, AI656081, H43421, N74128, AA612838,
			to 1390, where both a and b	9, AW021549, AA8113
			correspond to the positions of	AI862684, D53036, AA677516, AI382574, M79140,
			nucleotide residues shown in SEQ ID	F03274, AA356468, N89554, R7
			NO:76, and where b is greater than	N51175, AC005003, AL096880, A81324, A81326,
			or equal to a + 14.	AF181071
77	HBIBV81	665154	Preferably excluded from the	, AW302605,
			present invention are one or more	AA176780,
			polynucleotides comprising a	AW008216, AI949793, AI570129
			nucleotide sequence described by	AA025230, W37867, AI564486, AI252216, AI499109,
			the general formula of a-b, where a	AW242964, AI480359, AA774674, AI026080,
			is any integer between 1 to 768 of	_
			SEQ ID NO:77, b is an integer of 15	AW169773, AI350790, AA960795, AA300132,
			to 782, where both a and b	AA134208, AI695964, AI824322, AI377546,

				6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6
			correspond to the positions of nucleotide residues shown in SEQ ID	A168/600, AA468/88, A13800/8, AA91031/, A1061219, A1915290, A1277301, AA887251,
			Z.	, AA353348, AA419130, AA937303,
			or equal to a + 14.	AI025658, AA864265, H17773, AA931728, R50817, R27626, AA419069, AA897064, H17656, AI580298,
				AI521898, AW086126, AA626445, AI918844, F10191, F10115, T63745, R88235, AA721159
78	HSXBP02	062999	Preferably excluded from the	W24040, N95428
			present invention are one or more	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 264 of	
			SEQ ID NO:78, b is an integer of 15	
			to 278, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:78, and where b is greater than	
			or equal to a + 14.	
79	НСОСО19	668040	Preferably excluded from the	AA837754, AA581115, AC004466
			present invention are one or more	
			polynucleotides comprising a	
			nucleotide sequence described by	
			is any integer between 1 to 814 of	
			SEQ ID NO:79, b is an integer of 15	
			to 828, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:79, and where b is greater than	
			or equal to a + 14.	
80	HHENT19	985899	Preferably excluded from the	AC004998, AF130343
			present invention are one or more	
			polynucleotides comprising a	
			the general formula of a-b, where a	

			is any integer between 1 to 328 of SEQ ID NO:80, b is an integer of 15 to 342, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:80, and where b is greater than or equal to a + 14.	
81	HMTMB52	668717	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 523 of SEQ ID NO:81, b is an integer of 15 to 537, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:81, and where b is greater than or equal to a + 14.	AA031331, AA447922, AA878870, AA625391, AA026657, AI476276, AA305075, AA148792, AA446846, AW169122, AW149768, AI796276, AA603456, AA090696, AI566470, AA026887, AA455761, AA046950, AA837404, AW196971, AI636657, AA279066, AA321648, AA046476, AI025283, T30865, N40879, AA446847, AA845528, AI879232, AA188287, AA403246, H90077, AA128964, AA031332, H78109, N78226, T36197, AA936074, H37884, AA256024, T34431, T34451, AA308443, W51863, AA568448, AA877372, AI358381, AI351514, AA030022, H27053, AL048514, AF110777, AF151895, AF195950, U21858, I23471
83	HOGAL 19	668753	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 278 of SEQ ID NO:82, b is an integer of 15 to 292, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:82, and where b is greater than or equal to a + 14.	AA916322, AA244285, R57426, T67759, AF161472, AL117545, AL117608
83	HCQAG50	671361	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by	W27182, W31603, W76194, A1057607, AA203214, N43872, AI719074, AW009568, AW418865, R23692, AA136421, H27226, AA055206, AW372559, AW372556, AW372575, AW372567, AW195824, AI057563,

			the general formula of a-b, where a	AW372571, I95748	
			iny integer between 1 to 338 of		
			SEQ ID NO:83, b is an integer of 15		
			to 352, where both a and b		
			correspond to the positions of		
			nucleotide residues shown in SEQ ID		
			NO:83, and where b is greater than		
			or equal to a + 14.		
84	HDPLC22	674203	Preferably excluded from the	AI936345, AA887285, AI678802, AI41818	.87,
			present invention are one or more	AW058484, AI978830, AA614465, AI63478	184,
			polynucleotides comprising a	AA508486, AW299537, AI361913, AW299894	194,
			nucleotide sequence described by	AI359657, AI659077, AI718694, AI686164	.64,
			the general formula of a-b, where a	AA533665, AI469389, T90037, AA878577,	', AI739343,
			is any integer between 1 to 390 of	AA903695, AI934233, AI444978, AA910595	35, C00551
			SEQ ID NO:84, b is an integer of 15		
			to 404, where both a and b		
			correspond to the positions of		
			nucleotide residues shown in SEQ ID		
			NO:84, and where b is greater than		
			or equal to a + 14.		
85	HBMX090	674745	Preferably excluded from the	AI819368, AW043804, AA995625, AA15825	155,
			present invention are one or more	AI989534, AI632079, AI632177, AW385262	62,
			polynucleotides comprising a	AI436651, AA579669, W92990, AA58308	AA583087, AA702208,
			nucleotide sequence described by	AI767610, AI022894, AI694633, AI687149	.49,
	•		the general formula of a-b, where a	AA811023, AW207705, AA043498, AA062551	51,
			en 1 to 1541	AA024830, AA043234, AA024786, AI636005	, 500
			SEQ ID NO:85, b is an integer of 15	AA063156, AI564317, N99809, AI023039, W93177	), W93177,
			to 1555, where both a and b	AA583864, AA210642, AW388581, AI630114	114,
			correspond to the positions of	AW352131, T95281, AW388505, AW016381,	l, AW376106,
			nucleotide residues shown in SEQ ID	AW376098, AW082627, AA074329, AW339405	105,
			NO:85, and where b is greater than	AI479095, AA215346, AB011098, Y08686,	5, AF004830,
			or equal to a + 14.	U27455, X95642, U15555, AF111168	
98	HLMIS22	674761	Preferably excluded from the	AA320525, AW025411, AI653685, AI684617	517
			present invention are one or more		
			polynucleotides comprising a		
			nucleotide sequence described by		

			the general formula of a-b, where a is any integer between 1 to 441 of	
			SEQ ID NO:86, b is an integer of 15	
			to 455, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:86, and where b is greater than	
			or equal to a + 14.	
87	HE8AG73	677212	Preferably excluded from the	AA287561, AA307431, AA331618, AA287878, AF136450
			present invention are one or more	
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 661 of	
			SEQ ID NO:87, b is an integer of 15	
			to 675, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:87, and where b is greater than	
88	HCYBF14	683259	Preferably excluded from the	AA305210, AW015627, AI580368, AI830042,
			present invention are one or more	AI769572, AI741672, AA854575, AI073885,
			polynucleotides comprising a	AA834403, AA962811, AA757628, AI923528, N32611,
			nucleotide sequence described by	R79828, AW241940, AA961638
	-		the general formula of a-b, where a	
			is any integer between 1 to 479 of	
			SEQ ID NO:88, b is an integer of 15	
			to 493, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:88, and where b is greater than	
			or equal to a + 14.	
68	HKAAS37	682895	Preferably excluded from the	
			present invention are one or more	
	-		polynucleotides comprising a	
			nucleotide sequence described by	and the second s

			the general formula of a-b, where a is any integer between 1 to 402 of	
			to 416, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:89, and where b is greater than	
			or equal to a + 14.	
90	HBXFP72	688040	Preferably excluded from the	AA039596, AI871723,
			present invention are one or more	AI201016, AW338088, AI923596, AW268550,
			polynucleotides comprising a	AA826433, AA165179,
			nucleotide sequence described by	
			the general formula of a-b, where a	AA526606, R80933, AW131872, AA367044, AA278895,
			1 to 14	AI352299, AI688836
			SEQ ID NO:90, b is an integer of 15	
			to 1467, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:90, and where b is greater than	
			or equal to a + 14.	
91	HFIYP15	688044	Preferably excluded from the	N71063, W94295, N66037, H63245, AI568915,
			present invention are one or more	N73616, AA773642, W02603, AA907572
			polynucleotides comprising a	
			nucleotide sequence described by	
			a-b, where	
			is any integer between 1 to 1779 of	
			to 1793, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:91, and where b is greater than	
			or equal to a + 14.	
92	HEBAG86	688077	Preferably excluded from the	AA062994,
			present invention are one or more	I127084, AA406051,
			polynucleotides comprising a	AA978358, AI347575, AA405217, AA764978,
			nucleotide sequence described by	AI375843, W90774, AA649214, AA824648, AA258212,

			the general formula of a-b, where a	AI190424, AI719533, AI277627, W24232, AI095177,
			is any integer between 1 to 524 of	AA926760, AA927352, AA909449, AA284241, W56767,
			SEQ ID NO:92, b is an integer of 15	342704, H01228, AA514
			to 538, where both a and b	AA905893, F36912, D55269, H00845, R95095,
			correspond to the positions of	AI240810,
			nucleotide residues shown in SEQ ID	AA468124, D54053, AI
			NO:92, and where b is greater than	AA947003, T36306, AI597616, AA729251, W90739,
			oa + 14.	F33384, F28057, AI370811, R16001, AI144512,
				AA988588, AA033634, AA034341, R16102, AA826454,
				F27110, F31119, D52954, AA631119, AA894720,
				AI368559, F29378, AA489478, AA483450, F36024,
				D54336, T31368, T79165, C00350, AA594514,
				, N76784, D57671
				D59289, F00481, F32076, Z24770, AA301716,
				7, AW301847, F25224
			-	
93	HLDNM81	691124	Preferably excluded from the	AA345522, AI908286, AA484151, C20958, AA913510,
			present invention are one or more	AI678105
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			Ď	
			SEQ ID NO:93, b is an integer of 15	
		.=	to 483, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
	,		NO:93, and where b is greater than	
			or equal to a + 14.	
94	HARNC71	691721	Preferably excluded from the	H92130, AA468843, AA468855, AW244043, AA935265,
			present invention are one or more	F13685
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 705 of	
			SEQ ID NO:94, b is an integer of 15	
			to 719, where both a and b	

								-
			d to the po					
·			nucleotide residues shown in SEQ ID					
			NO:94, and where b is greater than					
			or equal to a + 14.			-		
95	HE20C31	693582	Preferably excluded from the	AA425207,	AA328348,	AI422986,	AW085230,	• •
			present invention are one or more	AJ223956,	X86000, Y09488	)9488		
			ψ					
			nucleotide sequence described by					
			the general formula of a-b, where a					
			is any integer between 1 to 599 of					
			SEQ ID NO:95, b is an integer of 15					
								-
			correspond to the positions of					
			nucleotide residues shown in SEO ID					
			NO.95 and where b is greater than					
			1 2 - 1					
			or equal to a + 14.			- [		T
96	HTXKQ20	696007	Preferably excluded from the	AA167034,	AA167085,		AA114130,	
			present invention are one or more	AI128866,	AA291957,	_	AA723387,	
			polynucleotides comprising a	AA766022,	AA299755,	AA991579, AA2	AA291956,	
			nicleotide semience described by	AW160699.	AI083889,	H38599, AA811	AA811428, AI348079,	9
			The more of the state of the st	יאסקנאגא	00202044		D34159 AA009919	
			WILCLE	יוויייייייי	1011111111	TECH LOCATOR	10100000 100	
			is any integer between 1 to 802 of	AI829155,	AA010462,		AI674507,	
			SEQ ID NO:96, b is an integer of 15	AI082075,	AA039391,	AC005041, ALC	AL049296	
			to 816, where both a and b					
			correspond to the positions of					
			nucleotide residues shown in SEQ ID					
			NO:96, and where b is greater than					
			or equal to a + 14.					
- 62	HE20K20	697955	Preferably excluded from the	AA328692,	AW244141,	AI435184, AC	AC005084, AC003093	093
			present invention are one or more					
			polynucleotides comprising a					
			nucleotide sequence described by					
			the general formula of a-b, where a					
			is any integer between 1 to 563 of					
			SEQ ID NO:97, b is an integer of 15					
			to 577, where both a and b					

			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:97, and where b is greater than	
			or equal to a + 14.	
86	HMWIW31	890869	Preferably excluded from the	W01234, AP000500
			present invention are one or more	
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 470 of	
			SEQ ID NO:98, b is an integer of 15	
			to 484, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:98, and where b is greater than	
66	HCEEH33	702853		AI436583
			present invention are one or more	
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 427 of	
			SEQ ID NO:99, b is an integer of 15	
			to 441, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
-			NO:99, and where b is greater than	
			or equal to a + 14.	
100	HAGBL85	703700	Preferably excluded from the	AW138272, AA976107, AA781938,
			present invention are one or more	AI093184, AA535789, AI803509, AA412322, AI216808
			polynucleotides comprising a	
•			nucleotide sequence described by	
			a-b,	
			is any integer between 1 to 510 of	
			SEQ ID NO:100, b is an integer of	
			e both a	

			correspond to the positions of	
			$\sigma$	
			or equal to a + 14.	
101	HLWAY38	705461	Preferably excluded from the	AA283147, AA142881,
			present invention are one or more	, AA23336
			polynucleotides comprising a	AI739455, AC007785
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 600 of	
			SEQ ID NO:101, b is an integer of	
			15 to 614, where both a and b	
			correspond to the positions of	
			NO:101, and where b is greater than	
			l to a + 14.	
102	H2LAN34	705692	Preferably excluded from the	AA314140, AI623759, AA768701, AW005593,
			present invention are one or more	AI420537, H65282, D58283, D80043, D57483,
			polynucleotides comprising a	D80253, D80164, D80366, D80022, C14331, D59467,
			nucleotide sequence described by	D59859, D81030, D80391, D59787,
			the general formula of a-b, where a	D80195, D59275, D59619, D80210, D51799, D80240,
				5, D80227, D80188, D80024, D59502,
			SEQ ID NO:102, b is an integer of	5, D59889, D80212, C14389,
				ω,
			correspond to the positions of	), AA305409, D80241, D80045,
				D51060,
			NO:102, and where b is greater than	2, AW179328, D80134,
			l to a + 14.	
			1	, D51250, AW377671,
				), D80168,
				C14227, AW178762, AW360811, AA514188, C14298,
				D81111, D80064, AW177501, AI910186, AW177511,
				C14407, Z21582, AA514186, D80133, AW177505,
				AI905856, C05695, D80247, AW352117, D80132,
				AW176467, AW375405, AW378540, D80268, AW179024,
				AW366296, AW360844, AW360817, AW375406,

			AW378534, AW179332, AW377672, AW179023,
			78905, AA285331, AI557751, AW36083
			2170, D80302, AW352171, T11417, D804
			5, AW178906, AW177731
			, D59373, AW179020,
			AW178909, AW177456, AW178980, AW179329,
			, AW179220, AW
			, AW179012
			25, AW367967,
			AW177728, AW179009, D51759, AW178774, AW178911,
			C06015,
			, T48593, D58101, D45260, T0311
			, AW177723, AA8091
			16, AJ1
			Y17188,
			78862, D34614, AF05869
			AR008278, X82626, I82448
			7, Y12724, AB012117
			B002449, A85396, AR066482, AR0
			, I19525, A86792
			$^{\circ}$
			514, AR060138, A45456, A26
			4, AR054175, AR066490,
			AR038669, AR066487, I18367, A3043
			187, AF13
			7, AR008281
			A70867, AR016691, P
			A64136, A68321, AR06013
			I79511, AR064240, U87247, AB023656, U79457,
			3, AR032065
			AR008382
103 HBMXT67	706204	Preferably excluded from the	4, AI918088, AI392950,
		present invention are one or more	, AI264604, AW304477,
		polynucleotides comprising a	356850,
		nucleotide sequence described by	AI383508, AI523998, AI276615, AA884388,

			the general formula of a-b, where a	AI356867, AI378377, AI589019, AI361250,
			between 1 to 187	
			SEQ ID NO:103, b is an integer of	AI766245, AA557471, AA632253, W95296, N68920, AI985972, R53701, W95553, AA736984, AI807975,
				W128999, H06890, AA470666, R44101,
			residues sho	
			allu wilete b is greater   to a + 14.	, T85379, R07731, M9
				F05911, AA325154, AA644479, T85478, T92165,
				F05899, R70373, N51810, AA091065, T93592, AW103327 AA936051 AA876718 H06849 1776421
104	HE2IE28	707161	Preferably excluded from the	H25350, H28544, AI955873, N29938, R12730,
			present invention are one or more	AL120665, AW104398, AC000064, AC007566
			polynucleotides comprising a	
			nucleotide sequence described by	
		-	the general formula of a-b, where a	
			is any integer between 1 to 239 of	
			SEQ ID NO:104, b is an integer of	
			15 to 253, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:104, and where b is greater than	
			or equal to a + 14.	
105	HBXCG73	707464	Preferably excluded from the	
			present invention are one or more	AW068217
			u	
				AA385863,
				U87166, AF176784, U17698
			is any integer between 1 to 691 of	
			105, b is an i	
			15 to 705, where both a and b	
			correspond to the positions of	
			ide residues	
			NO:105, and where b is greater than	
			or equal to a + 14.	

106	HATAN68	709015	Preferably excluded from the	AI796967, AW195747, AI740869, AA191594,
			present invention are one or more	AI498701, N64011, W86388, AI948435, AI125704,
				, D6100
			nucleotide sequence described by	AW204689, AI003139, AA2196
			the general formula of a-b, where a	346169, R20553,
			en	, T30719, A1197904
			SEQ ID NO:106, b is an integer of	5187, T24891, AI864073,
				C00222, AW020592, AW
			correspond to the positions of	, AI263584,
			residue	AW023863, AW021178, AW022826, AI538564,
			NO:106, and where b is greater than	AW019988, AI522052, AW023351, AW020931,
			or equal to a + 14.	AW020328, AI636727, AI871660, AW004606,
				AI884318, AW022308, AI579901, AW152182, N25033,
				AW020425, AW021693, AW022299, AI557238,
				AI633125, AW022981, AW020406, AW021182,
				, AA282824, AI935799,
				AI909697, AI887381, AI541321, AW023469,
				AI283760, AI744268, AI524626, AI571439;
				AW023884, AI915291, AW020629, AI696714,
				, AW022593, AI473536
				AI812091, W45039, AW020480, AW021466, AI359744,
				AI254731, AW020710, AI432030, AW198090, T69241,
				AI674423, AI932966, N21402, AI810544, AI889189,
				ò
				AW020295, AI925502, AW080157, AI866469,
				<u>,</u>
				AI625256, W74529, AA872507, AI368691, AW105296,
				AW02216
				AI699823, AW021717, AI539545, AA554929,
				103, AW022760, AI61327
		_		AW022874, AI469262, AW021930, AI421903,
				32, AI818980, U0694
				AL117587, A77033, A77035, L35261, A52184,
				AL137533, A44314, Z13966, AF200464, AF060555,
				AF124396, I32738

			recent intention are one or more	AA454978, A	AA132519,	AW135352,	AA179230,	N45125
_			present invention are one or more polynucleotides comprising a		AM004324, AA857184,	AI985060,	AI361206,	R54585,
			nucleotide sequence described by	AI272727, A	AI766581,	AI940540,	AA923780,	Z38507,
			the general formula of a-b, where a	AW192986, A	AC004685			
			is any integer between 1 to 452 of					
			7					
			15 to 466, where both a and b					
			correspond to the positions of					
			nucleotide residues shown in SEQ ID					
			NO:107, and where b is greater than					
			or equal to a + 14.			i		
108	HBJF165	711769	Preferably excluded from the	AI740525, P	AI800754,	AI189295,	AA878902,	
			present invention are one or more	AI262709, P	AW169159,	AA495986,	AI469879	
			polynucleotides comprising a					
			nucleotide sequence described by					
			the general formula of a-b, where a					
			is any integer between 1 to 309 of					
			SEQ ID NO:108, b is an integer of					
			15 to 323, where both a and b					
			correspond to the positions of					
			nucleotide residues shown in SEQ ID					
			NO:108, and where b is greater than					
			or equal to a + 14.					
109	HSNAL84	711840	Preferably excluded from the	н	AI339840, AV	AW044507, A	1216527,	AI620878,
			present invention are one or more		\1292180,	AI292180, AI358083,	AI954691,	
			polynucleotides comprising a		AA321122,	AA321123,	AA321122, AA321123, AI654341,	N46790,
			nucleotide sequence described by	AI953114, N	N69895, A.	N69895, AI970523, AA226346,	A226346, A	AA226347,
			the general formula of a-b, where a	AF196969				
			is any integer between 1 to 434 of					
			SEQ ID NO:109, b is an integer of					
			15 to 448, where both a and b					
			correspond to the positions of					
-			nucleotide residues shown in SEQ ID					
			NO:109, and where b is greater than					
,			or equal to a + 14.					

		519,			514,												953,		8834,							757,		5118,					503,		
AI79252	A1/55214	<#	AW438542,	AI133552,	T74524, AI627614	AW328202,	AI251203,	AA700943,	AA501461,	AI613389,	AI251284,	AI732483,	AL119247,	AA593537,	AI282253,	AA704393,	AA533025, H0795	AI473671,	AI040051, AA828	AI754767,	AA492495,	AW243793,	AI962030,	AI300054,	AA595661,	R94326, AA747757	AA687730,	AA530958, T05	AI267356,	AA814503,	AA319233,	AC006211,	Z84486, AP000503,	AL049643,	AC007421
AL042373,	~ ~	4,	AW304580,	AI114733,	AI380617,	AA831638,	AA916430,	AW068596,	AA456937,	AW303098,	AL042670,	AA833896,	AI223626,	AA618316,	AW069412,	AA483606,	AA719564,	$\overline{}$	R99034, A	AW089016,	AA524616,	AI254770,	AA603413,	AA315361,	AW020150,	AI499954,	AA581903,	AW271904,	AW275432,	AA632993,	AI003626,	AC006239,	AF134726,	AC007565,	AT,031228
Oι	260	5	AI792575,	AL079734,	AA503298,	AW328331,	AI053688,	AA600202,	AI284543,	AI859438,	AL042667,	AA833875,	AW069227,	AA487226,	AI923052,	AI278972,	AA502991,	AI859946,	AW023111,	AA809546,	AW026305,	AA534064,	AA584360,	AL041375,	AA568204,	AA013168,	AW408767,	AA630854,	AW238484,	AI251576,	AI362442,	AI537995,	AL109627,	AL096791,	AT.117694
AA442729,	AW23/905,	5410	AI440117,	AI335387,	AA904211,	AI521525,	AI754170,	AI251034,	AA526542,	AW270385,	AA714110,	AI890324,	AA536040,	AA524229,	AW084445,	AA535216,	AI687343,	AI799607,	AI817658,	AI081147,	AI733856,	AI309059,	AI254779,	539	AA659832,	AA019973,	AW089625,	AA410788,	AI369580,	AI745151,	AI583142,	AI291439,	AL031602,	AP000117,	AC004771.
excluded from the	rentic	polynucleotides comprising a	nucleotide sequence described by	the general formula of a-b, where a	ger between	SEQ ID NO:110, b is an integer of	wher	correspond to the positions of		NO:110, and where b is greater than	a + 14.																								
711878																																		-	_
HCRND41																-																			
110																																			

AC002302, AP000193, AF196779, AC004975,
663, AC004796, AC007052, AC00587
F134471, Y14768, AC005828, AC007664,
0501, U91323, AL109801
0, Z98884, AC006511, AI
0505, AL096701, AC0049
, AL049539,
, AC005037, AC00611
68, AC004653, AP00008
, AC005231, AC009247, AL02214
AL109628, AC005225, AC005599, AC002316,
AL022326, AL133448, AC006388, AC008372,
, AC005046,
, AC005971, AC007546, AC00566
712, AL049872, AL035086, AC00419
AL009181, AP000547, AL078477, AC004686,
AF045555, AB000876, AL034402, AC002553,
AL022069, AC010582, AL109798, AL031587,
AL035413, AC004531, AP000497, AC003041,
82, AC004156, AC003982,
19, AC007277, AL021154
9, Z94056, AC006450, AF067844,
_
882, AL034417, AP001053, U91318
, Z97184, AC004750, AC
, Z98946, AC005821, AC
, AL031283,
0, AC006312
AC002996, AP001052, Z82208, AC012627, AC004596,
4, AC005730, AC004804,
 AC007842, AC005535, AC006120, AP000510,
, AP000359, AC005280, AC00235
7934, AL122023, AC005538, AF111167,
L035407, AC005924, AC005777, AL0497
Z98200, AC006538, AC006530, U96629, AL117329,

				טייייייייייייייייייייייייייייייייייייי
				AFUUU694, ACUUSUBI, AB
		-		, AC002352,
				AC005071, AC007036, AC004812, AL034549,
				, AL021155
				AL031005, Z77249, AC005881, U95742, Z93241,
				', AC004106
				AC009509, AP000114, AL031659, AL121603,
				AC005291, AC007993, AC005666, AL023553,
				AL034548, AC008044, L78833, AC005183, AC004002,
				AC005529, AC005013, AC005300, AC006441,
				AC004560, AL049636, AL049779, AP000211,
				AP000133, Z83844, AL035400, AL031289, AC005039,
				6
				AC004821,
				6, AP000046, AL132992
				AC005030, AC007406, AC005015, Z99716, Z98751,
				AC004466, AC004874, AC004966, AC005095,
				AC005102, AC005245, AC002378, N50319, AA236194,
				AA236183
==	HPXAA41	712638	Preferably excluded from the	AA910497, AI701451, AA429326, AI743089,
			present invention are one or more	
			polynucleotides comprising a	AA315932, H51256, AA425105, AA921331, AA885637,
			nucleotide sequence described by	, AI680678
		٠	the general formula of a-b, where a	AW014324, W01644, AI809584, AI762128, AI693885,
			ger between	, AI457315,
			SEQ ID NO:111, b is an integer of	AI630807, AI694045, AA928976, AA931651, N71630,
			15 to 876, where both a and b	AW295247, AA094470, AF147430, D10920
			correspond to the positions of	
	,		nucleotide residues shown in SEQ ID	
			NO:111, and where b is greater than	
			or equal to a + 14.	
112	HHSFO42	713301	Preferably excluded from the	F36273, AA654968, AL119691, AI281881, AA515224,
			present invention are one or more	AW236342, AI358343, AI35169
			polynucleotides comprising a	AW079241, AA084070,
			nucleotide sequence described by	AI654247, AW304805, AI920876, AI688846,
			the general formula of a-b, where a	AI358813, AW168342, AA661948, AW020992,

	is any integer between 1 to 368 of	AI284640, AI192631, AA502103, AA347927,
_	112, b is an inte	AW276435, AI857789, AI291124, AW193265,
	82, where both a	AW270382, AA347930, AI567674, AW089322, T08638,
	to the positions	AI500454, AA515051, AW265385, T07451, AW089789,
	residue	5330, AI43130
	NO:112, and where b is greater than	AA593247, AI625244, AA284179, AI061313,
	oa + 14.	AL043721, AI858451, AA829106, AW029038,
		AI962050, AI291268, AI339850, AW238278,
		AI350211, AA555229, AA324849, AA720025,
		AA594145, AW193432, AW440836, AI434311,
		AI
		AI133636, T06828, AA528516, H86305, AA513999,
		9, AA491767, AI1512
		, AA225
		, AA747472,
		81697,
		, AI355224, N95
		AA847499,
		, AI061296, AI358571,
		2, AI674873, AI
		_
		85124, AA719805, AA349366, AW33956
		, AW166815,
		, H63607, AI
		AA483731, AW302450, AA782318, AA826303, N75391,
		, AI049722, AW152057, AI672135, R83
		ΑI
		, AW021583, F37169,
		AA594725, AI571562,
		104375
		2, AI017024, AA782272, AA0741:
-		, AA857486, AI801591
		U51696, AI148245, R95171, AA338522, AA634227,

2, H54443, AI
, AA584201, AA78911
, AL046409, AI824562,
298, AW008952, AW438643, AA67967
AA371533, AA385100, AI721122, AL049829, Y10196,
M37551, AL109984, U14719, AP000553, U14718,
U14711
AP000032, U14714
629, S77605, 1
X55926, X75335,
, AC016026, U14705
 AC005516, U47924, Z73358, A3997
020, AC004821, U57009, AP
993, AC006057, AC005081,
, AC
, AL122020, AL049758,
AC004886, AF1904
X55925, AC005747, AL10
 U57006, U18394,
707, AC00
AC004907, AC004034, U02057, AC004592, U14700,
. AC005939, U14710, AL049779,
1846, AC004955
U18387, AL034549, AC008101, Z15025, AC004651,
555, AC007227, AC006111, AF2
381,
48, U57008, AC006538, X55932, AC
2318, U02063, AC016830, AF020503,
 AC007387
AL132987, AL050318, U73024, AC004239, U14702,
4685, U14687, U1
77058, AC006039, AC005288, AL023574
84469, AC004876, AC005215, AC005866,
AC005856, AP000497, AC004087,
 031677, AP000133, AP000211, Z8
U49740, M19045, J03801, E01888, E02193,

				אפאפק פכוחחתפה דחכחחתה וחדאפתוה אוכחחתפה
				ADOJOTA, AFOOCATA, EJOAR
				4, ACUISUZ/, ACUUSIJS, ACUU4SSI, ASSUS
				1, AC005846,
				AC005041, AP000228, AC010072, AP000255, X54178,
				AE000658, AC004019, U12582, AL035464, AC005225,
				AF165147, U14686, U14688, U14689, AP000047,
				AC007687, Z82976, AL109628, AC008064, AC004453,
				AC004235, AL080316, AP000140, AC007444,
				$\circ$
				AC007488, AL024498, AC005166, AP000031,
				AL079340, AC005480, AL008721, AL033376,
				AC005696, AL117258, U18400, AL031427, AC007157,
				AC007971, AD000092, AC006160, AF064863,
				AC005035, AC009498, AC005104, AL031985, U62317,
				AL133485, Z84721, X54181, AF070718, AL035071,
				AL023575
113	HCEIE94	714156	Preferably excluded from the	H16630, Z46007, R18668, AA779244, AI742776,
			present invention are one or more	AA648586, AL049824
_			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
_			SEQ ID NO:113, b is an integer of	
			re both	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:113, and where b is greater than	
			or equal to a + 14.	
114	HWLQA43	714877	Preferably excluded from the	
		_	present invention are one or more	
		_	polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	

			is any integer between 1 to 357 of SEQ ID NO:114, b is an integer of 15 to 371, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:114, and where b is greater than or equal to a + 14.	
	HFXHM92	715343	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 567 of SEQ ID NO:115, b is an integer of 15 to 581, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:115, and where b is greater than or equal to a + 14.	AA628088, AI640353, AI767467, AI921798, AI810416, AI810681, AI125878, AW073826, AA182773, AI954699, AW293730, AI828389, AI580952, AI313461, AI651050, R49049, AA772043, N33101, AA496896, AA928502, AI031772, AA864471, AI401678, W45009, N93870, T63242, AW194759, AI291264, AI983163, H99107, AI038482, AA604895, AI292264, AI983163, H99107, AI038482, AA604895, AI292264, AI983163, H99107, AI038482, AA604895, AI590061, AI452404, AA099425, AI200622, AA974079, R73465, AI432100, AA081421, AI698923, AI571941, AA480265, AI017448, N64632, AI199075, AA58942, AI982600, T33000, H65188, AI817275, AA584978, N55816, AI827372, AI701353, H28606, AA5947626, AA722706, H00447, AI826589, N24348, R55743, H03410, Z41694, F02896, R62185, H45540, N92338, N78793, F26109, N48260, N44922, W38368, AW003189, H00490, AI310166, AI370047, AI313291, AI340484, T90688, R64574, N93394, H45548, F35733, AI308272, AA320558, AA365588, W24522, AA651757, N45701, AA096344, AA099538, N50614,
116	HHSGE44	716212	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 691 of SEQ ID NO:116, b is an integer of	W61017, W61009, AA128255, AA806126, AA136365, AA936371, AW150765, AI279977, AA125840, AW448960, AA128312, AA125856, AI186377, AI269647, AI265821, AA873528, AA136280, R07851, R62482, AB031039, AB031040

			15 to 705, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			ಹ	
			or equal to a + 14.	
117	HWLQ133	717222	Preferably excluded from the	, AI935263, AI741826,
			present invention are one or more	AI809826, AI524082, AW102619, AI804122,
			polynucleotides comprising a	AI540191, AW405122, AI589161, AI860317,
			nucleotide sequence described by	AI742839, AA631117, AI809767, AI363498,
			a	AA427510, AA480840, AI298899, AI086078,
			teger between	AA847830, AI354791, AI051661, AI675663,
			SEQ ID NO:117, b is an integer of	AI122706, AI37777, AA503878, AI202112,
			6, where both	AI720306, AI394378, AI375978, AA764814,
			correspond to the positions of	AW406420, AA333282, AI286081, AI298712,
			nucleotide residues shown in SEQ ID	AA292572, AI492636, AI867099, AI417659,
			NO:117, and where b is greater than	AI468559, AA971791, AI222922, AA913022, H96909,
			or equal to a + 14.	AA972549, AA992120, AI928774, AW207223,
				AA814866, AA757376, N35604, AA907092, AW072106,
				AW166047, AA757394, AI345182, AA457729,
				AA235704, AW183289, AA430669, Z57528
118	HFIAW90	718259	Preferably excluded from the	H19365, AA286719, N44262, AA447218, AA974582,
			present invention are one or more	AC004817
			polynucleotides comprising a	
	-		nucleotide sequence described by	
			the general formula of a-b, where a	
			en	
			SEQ ID NO:118, b is an integer of	
			15 to 975, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:118, and where b is greater than	
			or equal to a + 14.	
119	HOSEP43	719829	Preferably excluded from the	
			present invention are one or more	
			polynucleotides comprising a	
			nucleotide sequence described by	

			the general formula of a-b, where a	The second secon
			teger between	
			SEQ ID NO:119, b is an integer of	
			., where k	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
	_		and wh	
			or equal to a + 14.	
120	HUSGY48	721985	Preferably excluded from the	796, AA528524, AA886535,
			present invention are one or more	Ä
			polynucleotides comprising a	AI914394, AI392630, AI338999, AI123694,
	-		nucleotide sequence described by	AA977864, N64163, AA359165, AA975366, AI126532,
				AA916014, AI832970, AI351443, AA888167,
			en	AI093884, AA973611, AI833308, F33502, AW264528,
			SEQ ID NO:120, b is an integer of	, AI039168,
			, where both a	AA340839, AI814033, AI682171, AI719390, F32392,
			correspond to the positions of	AI268514, F103
			residues shown in	R02355, AI742966,
			NO:120, and where b is greater than	R38577, Z20180, D19917, AW419258, AI963625,
			l to a + 14.	5, AI345612, AI34
				AL037602, AI590043, AI699011, AI364788,
				AI623535, AI064830, AW161156, AL079728,
				4, AL036403,
				, AI358701, AL110306,
				0, AL121
				AA748353, AW082113, AI866465, F30529, AI500061,
				AW440167, AL045500, AL119399, AI269862,
				AL046466, AI923989, AI249877, AW403717,
				AI582912, AI538885, AA641818, AL036631,
				AL038605, AL036638, AI567582, AW162194,
		_		AL043168, AW071417, AI698391, AW151136,
				AI538850, AI285439, AI582926, AI801793,
				AW087445, AI433157, AW020373, AI554821,
				AL121328, AI923509, AI539771, AI859991,
				AI537677, AL047763, AW021717, AI500659,

	AI815232, AI801325, AI500523, AI582932,
	I284517, AL043089, AI500706, AI49177
	I445237, AW151138, AI521560, AI88918
-	AI500662, AI284509, AI889168, AL120695,
	66573, AI633493, AI434256, AL11040
	38661, AI284513, AI888118, AL03745
	355779, AI44025
	7882, AI473536, AI43303
	, AL038529, AI3075
	AW009337, AI344817, AI859464, AI335208,
	, AW051088,
	AL119457, AI919593, AA572872, AW129264,
	81, AL119791, AI866469, AI34546
	, AW160916,
	, AI621341, AI824576, AI2419;
	AI539153, AI37125
-	, AL043981, AI698401, AW0838(
	7, AI690946, AL039
	63, AI969655, F35299, AI
	865, AI624293, AA768046,
	, AI683492, AL045163, AL036
	0, AL042745, W38553, AI89112
	42628, AI887775, AI872423, AW
	239, AI611743, AW020876, AI
	, AL042382, AI440263,
	242,
	, AL039086, AW162189, AI25472
	91710, AL043152, AW160905, AI34478
	197, AF04495
	AR043114, X64898, A74894, I33392, AL137529
	978, AL117648, A65341
	16, AL13306
	, A58545, AL049430, AL122100, AF
	U96683, A08916, A08910, A08909, AB01622

	AF126247, I89947, A08908, AF158248, A86558,   U87620, AL137521, AR038969, AR034821, AL137479
-	090943, AL133072, A08913, AL122049, AJ01275
	, AL133606, AL049382
	557, A08912, X06146,
	908, AL137463, I48979, AJ000937,
	AF078844, AR013797, I89931,
	133049, I49625
	938, AF031147, U5899
	AL122050, I42402, U49434,
	AL096751, U88966, I79595, AL122110, Y10823,
	AL133080, A83556, AL13745
	5, U72620, AR020905, AF113694, AE
	F028
	X83508, AR038854,
	48, Y16645, AF067728, XT
	95114, AF118070, AF065135, A77033, A7703
	3, A93350, AL137711,
	U35846, AF
	458, AL080124, AF111112, U67958,
	8, AL110218, A18777, AR068753, AL137
	6, AF061836, AL117583
	AL080127, E
	18, AL13303
	9466, AF111851, AF090886, AL1330
	332, AF113013, I00734, E07361, AF10
	ω
	AL137271, AL133081, AL050172, AF210052,
	AL137550, E00617, E0
	AL137547, AL137294, AB008792, E02221, S78214,
	U775
	Y10936, X62580, AL11
	4,
	AL050155, AL133619, L
	AF008439, AL122123, X72889, A58524, A58523,
	1 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

			, A76335, E02349, AF183393, Y14314,
	4.00		AL023657, AF090901, X93495, AL133565, AF113691,
			5. AL110158, AF090900, AL133014,
			4, ALOSO138, ALOSO393, E15569,
			17, AL133640, AL117585, AL1
			1, AL137658
			AF079765, A
			5, AF067790,
			AL050024, AF113699, AL137560
121 HSLEC18	722249	Preferably excluded from the	AA826487, AA339231, AI
		present invention are one or more	AL138182, AA502991, AW328000, AI815210,
		polynucleotides comprising a	
		nucleotide sequence described by	8, AI361090, AI573198,
		the general formula of a-b, where a	AA503298, AL038842, AI859438, AA812684,
		eger between 1	AW238253, AA714110, AI962030, AL042373,
		SEQ ID NO:121, b is an integer of	AI612142, AA578621, C06004, AI799569, AW341978,
		15 to 2043, where both a and b	_
		correspond to the positions of	AI049955, AA515728, AW327624, AI752365,
		residue	, AI025930,
		NO:121, and where b is greater than	, AI362442,
		or equal to a + 14.	AA598892, AI653776, AW274191, AA652834,
			5, AW410784,
			AA833875, AI
			AW082104, AI340832, AA302812, AA993636,
			, AW088631, AI306232,
			6, AI824476,
			N23504, AA228349, AW190484, AA595499, AL048275,
			AI207424, AA832444, AA548610, AL079734,
			AI583466, AA493226, AL118991, AI431513,
	-		AL037632, AA828047, AW081303, N41775, AI360558,
			AI275982, AL041375, AI821987, AI251576,
-			AI610737, AI732869, AL031602, AC005231,
			2, AL031311, AC005081, Z84
			6211, AC002316, AF196779, AL133371
			AL022723, AC007542, AL035414, U80017, AF030453,

5, U62293, ALO49776, ACO06323, ALO 8, ALO35684, ACO02429, ACO022996, 5, ALO34379, ALO49540, ACO023667, 0, ACO04821, Z82198, ACO05037, AC 7, ALO31774, AD000813, AC005512, 3, AC006547, AC004813, AF184110, 5, ALO31289, Z93023, AP000555, X5- 7, AC005484, ALO08637, ALO31005, 6, AC002484, ALO08637, ALO31005, 6, AC002484, ALO08637, ALO31005, 6, AC002484, ACO04659, AC005940, 8, AL135783, Z98036, AC005940, 8, AL135783, Z98036, AC005940, 9, AC012627, ALO35455, D88270, AC 1, ALO31984, AC002550, Z82214, AC 6, AC002663, AC002550, Z82214, AC 9, AL133246, AC007371, AL049757, 7, AC006071, AC004590, AL136295, 1, AC005881, AC004967, AF200465, 3, AC00581, AC004815, AF204665, 3, AC005881, AC004887, AL078634, 6, AC00581, AC004815, AC010077, 3, AC00581, AC004815, AC04019, 1, AC007878, AC004148, AC004019, 1, AC005664, AL049761, AC007993, 11, AC006661, AP000692, AC005011, 9, AC00661, AP0006312, AL022165,	AC00501	015, AC004914, AC004213, Z95115, AC002551,
B, ALO35684, ACO02429, ACO02996, ALO34379, ALO49540, ACO03667, O. ACO04821, Z82198, ACO05037, ACO ALO35174, AD000813, ACO02073, ALO31074, AD000813, ACO05512, ALO31289, Z93023, AP000555, X57, ACO05484, ALO36837, ALO31005, ACO0547, ACO3029, AP000555, X57, ACO12627, ALO31689, ACO0250, ACO05940, ACO12627, ALO31884, ACO04659, ACO05940, ACO12627, ALO31884, ACO02550, Z82214, ACO12627, ALO34545, D88270, ACO12627, ACO02550, Z82214, ACO12627, ACO02550, Z82214, ACO12627, ACO02550, Z82214, ACO134549, ACO05670, AF111169, ACO05683, ACO05670, AF111169, ACO05681, ACO05670, AF101169, ACO05881, ACO04967, AF200465, 3, ACO05881, ACO04967, AF200465, 3, ACO05881, ACO04967, AF200465, 3, ACO05881, ACO04815, ACO10077, 3, ACO08040, ACO04815, ACO04019, 1, ACO08788, ACO04815, ACO04019, 1, ACO0564, ALO49761, ACO07993, 1, AF038458, AD000692, ACO05011, ACO06681, AD000692, ACO05011, ACO06681, ALO32165, ACO06312, ALO22165,	AL121	5, U62293, AL049776, AC006323
ALO34379, ALO49540, ACO033667, ACO04821, Z82198, ACO05037, ACO ALO31774, AD000813, AC002073, ALO31774, AD000813, AC005512, AC006547, AC004813, AF184110. ALO31289, Z93023, AP000555, X5-ACO05484, ALO8637, ALO31005, ACO05484, ALO8637, ALO31005, ALO31984, AC004659, AC005940, ALI35783, Z98036, AC002310, ACO ALI35783, Z98036, AC002510, ACO ALI35783, AC00550, Z82214, ACO ALI35783, AC005670, AF111169, ACO03066, AC00572, ALO50404, ACO0506, AC00572, ALO50404, ACA ALI33246, AC007298, AC005548, AC005280, AP000967, AF200465, AC005881, AC004967, AF200465, AC005881, AC004590, ALI36295, ACA AC005519, AC004815, AC004816, AC004816, AC0049148, AC004914, ACA AC00564, ALO49761, AC007993, AC005641, AD00692, AC005011, AC006468, AC006312, AL022165, AC006468, AC006312, AL022165, AC006468, AC006312, AL022165,	AC005	8, AL035684, AC002429, AC0029
AC004821, Z82198, AC0055037, AC0 AL031774, AD000813, AC002073, AL079301, AL035587, AC005512, AC006547, AC004813, AF184110, AL031289, Z93023, AP000555, X55, AL031984, AC004659, AC005940, AL135783, Z98036, AC002310, AC, AC012627, AL035455, D88270, AC AC012627, AL035455, D88270, AC AC002477, AC002550, Z82214, AC AC003663, AC005670, AF1111169, AC003663, AC005670, AF1111169, AC005003, AC007371, AL049757, AC005003, AC007371, AL049757, AC005280, AP000967, D86995, AC AL133246, AC00259, AC005548, AC005881, AC004590, AL136295, AC005881, AC004590, AL136295, AC005881, AC004687, AL078634, AC005519, AC004687, AL078634, AC00564, AL049761, AC007993, AC00564, AL049761, AC007993, AC005661, AP000692, AC005011, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AP000	45, AL034379, AL049540, AC0036
AL031774, AD000813, AC002073, AL079301, AL035587, AC005512, AC006547, AC004813, AF184110, AL031289, Z93023, AP000555, X5, AC005484, AL008637, AL031005, AC005484, AC004659, AC005940, AL135783, Z98036, AC002310, AC, AC012627, AL035455, D88270, AC AL034549, AP000275, Z93241, D8, AC003663, AC005670, AF111169, AC003663, AC005670, AF111169, AC005280, AP000967, D86995, AC, AL133246, AC007371, AL049757, AC005280, AC004967, AF200465, AC005281, AC004967, AF200465, AC005881, AC004967, AF200465, AC005881, AC004590, AL136295, AC005881, AC004687, AL078634, AC005519, AC004687, AL078634, AC00564, AL049761, AC007993, AC00564, AL049761, AC007993, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AF207	50, AC004821, Z82198, AC005037
ALO79301, ALO35587, AC005512, AC006547, AC004813, AF184110, ALO31289, Z93023, AP000555, X5-AC005484, AL008637, AL031005, AC005484, AC004659, AC005940, AL031984, AC004659, AC002310, ACAC012627, AL035455, D88270, ACO12627, AL035455, D88270, ACO12627, AC002560, Z82214, ACAC002477, AC002550, Z82214, ACAC002603, AC005570, AF111169, AC005280, AC005572, AL0349757, AC005280, AC00552, ACAC005881, AC00258, ACO05581, AC005881, AC004597, AC005881, AC004597, AC005881, AC004596, AC010077, AC005611, AC004687, AL036295, ACAC005611, AC008040, AC004815, AC004919, AC008646, AL049761, AC00561, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165, AC006468, AC006312, AL022165,	AL020	, AL031774, AD000813,
AC006547, AC004813, AF184110, AL031289, Z93023, AP000555, X5- AC005484, AL008637, AL031005, AC005484, AC004659, AC00553, AL031984, AC004659, AC005940, AL135783, Z98036, AC002310, AC AC012627, AL035455, D88270, AC AC002477, AC002550, Z82214, AC AC002477, AC002550, Z82214, AC AL034549, AP000275, Z93241, D8 AC005603, AC005670, AF111169, AC005280, AC005722, AL050404, AC AL133246, AC005722, AL050404, AC AL031575, AC002580, AC005512, AC005881, AC004967, AE200465, AC005881, AC004560, AL136295, AC005881, AC004687, AL078634, AC005648, AC04487, AL078634, AC005644, AL049761, AC007993, AC00564, AL049761, AC007993, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AP000	, AL079301, AL035587, AC005
AL031289, Z93023, AP000555, X5- AC005484, AL008637, AL031005, AP000065, AC003029, AP000553, AL031984, AC004659, AC005940, AL135783, Z98036, AC002310, AC AC012627, AL035455, D88270, AC AC002477, AC002550, Z82214, AC AL034549, AP000275, Z93241, D8 AL034549, AP000275, Z93241, D8 AC005003, AC005721, AL049757, AC005003, AC005722, AL050404, AC AL133246, AC005722, AL050404, AC AL133246, AC005729, AC005412, AC005881, AC004967, AF200465, AC005881, AC004967, AF200465, AC005881, AC004580, AL136295, AC005881, AC004687, AL078634, AC005648, AC044815, AF064861, AC008040, AC004815, AF064861, AC00564, AL049761, AC007993, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AF139	3, AC006547, AC004813, AF184
AC005484, AL008637, AL031005, AP000065, AC003029, AP000553, AL031984, AC004659, AC005940, AL135783, Z98036, AC002310, AC AC012627, AL035455, D88270, AC AC002477, AC002550, Z82214, AC AL034549, AP000275, Z93241, D8 AL034549, AP000275, Z93241, D8 AC005003, AC005670, AF111169, AC007066, AC005722, AL049757, AC00508, AC005722, AL06944, AC AL133246, AC007298, AC005548, AC005881, AC004967, AF200465, AC005881, AC004967, AF200465, AC005881, AC004590, AL136295, AC00511, AC004687, AL078634, AC00511, AC004687, AL078634, AC008040, AC004687, AL078634, AC008040, AC004815, AF064861, AC008646, AL049761, AC004019, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AC003	5, AL031289, Z93023, AP000555, X
AP000065, AC003029, AP000553, AL031984, AC004659, AC005940, AL135783, Z98036, AC002310, AC, AC012627, AL035455, D88270, AC, AC002477, AC002550, Z82214, AC, AC003663, AC005670, AF111169, AC003663, AC005670, AF111169, AC007066, AC005722, AL050404, AC, AL133246, AC007298, AC005548, AC005280, AP000967, D86995, AC, AL133246, AC002116, AC005412, AC005881, AC004967, AF200465, AC005881, AC004967, AF200465, AC005881, AC004590, AL136295, AC005881, AC004687, AL078634, AC00519, AC004687, AL078634, AC008040, AC0048815, AF064861, AC008646, AL049761, AC007993, AF038458, AP000128, AP000206, AC006468, AC006312, AL022165,	AC004	, AC005484, AL008637,
AL031984, AC004659, AC005940, AL135783, Z98036, AC002310, AC AC012627, AL035455, D88270, AC AC002477, AC002550, Z82214, AC AL034549, AP000275, Z93241, D8 AC00503, AC005670, AF111169, AC00503, AC007271, AL049757, AC007066, AC007298, AC005548, AL133246, AC007298, AC005548, AC005280, AP000967, D86995, AC AL133246, AC004967, D86995, AC AL031575, AC002116, AC005412, AC005881, AC004967, AF200465, AC005881, AC004590, AL136295, AC005519, AC004590, AL136295, AC005519, AC004687, AL0786481, AC003071, AL096791, Z98752, Z8 AC00564, AL049761, AC007993, AC005664, AL049761, AC007993, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	ACO05	6, AP000065, AC003029, AP00055
AL135783, 298036, AC002310, AC, AC012627, AL035455, D88270, AC AC002477, AC002550, Z82214, AC, AL034549, AP000275, Z93241, D8, AC003663, AC005670, AF1111169, AC005003, AC00572, AL049757, AC005280, AP000967, D86995, AC, AL133246, AC005298, AC005280, AP000967, D86995, AC, AL031575, AC002116, AC005412, AC00581, AC004967, AF200465, AC006071, AC004590, AL136295, AC, AC005519, AC004591, Z98752, Z8, AC008040, AC004815, AF064861, AC00564, AL049761, AC007993, AC00564, AL049761, AC007993, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AL117	, AL031984, AC004659,
AC012627, AL035455, D88270, AC AC002477, AC002550, Z82214, AC AL034549, AP000275, Z93241, D8 AC003663, AC005670, AF1111169, AC005003, AC007371, AL049757, AL133246, AC007298, AC005548, AL031575, AC002116, AC005548, AC005280, AP000967, D86995, AC AL031575, AC002116, AC005412, AC005881, AC004967, AF200465, AC005881, AC004590, AL136295, AC00581, AC004687, AL078634, AC005519, AC004687, AL078634, AC005664, AL049761, AC004919, AC005664, AL049761, AC007993, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AC002	, AL135783, Z98036, AC002310
AC002477, AC002550, Z82214, AC AL034549, AP000275, Z93241, D8 AC003663, AC005670, AF111169, AC005003, AC007371, AL049757, AL133246, AC007298, AC005548, AL031575, AC002216, AC005548, AC005881, AC004967, B6995, AC AL031575, AC002116, AC005811, AC005811, AC004967, AF200465, AC005811, AC004967, AF200465, AC005811, AC004681, AL078634, AC005519, AC004687, AL078634, AC005644, AL049761, AC004019, AC005644, AL049761, AC007993, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AC006	530, AC012627, AL035455, D88270,
AL034549, AP000275, Z93241, D8 AC003663, AC005670, AF111169, AC005003, AC007371, AL049757, AC007066, AC005722, AL050404, AC AL133246, AC007298, AC005548, AC005280, AP000967, D86995, AC AL031575, AC002116, AC005412, AC005881, AC004967, AF200465, AC006071, AC004590, AL136295, AC005519, AC004687, AL078634, AC005519, AC004687, AL078634, AC008040, AC004815, AF064861, AC008640, AC04815, AF060931, AC005664, AL049761, AC007993, AC005661, AP000692, AC005011, AC006468, AC006312, AL022165,	AC005	, AC002477, AC002550, Z8221
AC003663, AC005670, AF111169, AC005003, AC007371, AL049757, AC007066, AC005722, AL050404, AC AL133246, AC007298, AC005548, AC005280, AP000967, D86995, AC AL031575, AC002116, AC005412, AC007358, AC004967, AF200465, AC00581, AC004590, AL136295, AC006071, AC004590, AL136295, AC005519, AC004687, AL078634, AC005519, AC004687, AL078634, AC005640, AC044148, AC004019, AC00564, AL049761, AC007993, AF038458, AP000128, AP000206, AC006468, AC006312, AL022165,	AC005	, AL034549, AP000275, Z93241, D
AC005003, AC007371, AL049757, AL0207066, AC005722, AL050404, AC AL133246, AC007298, AC005548, AC005280, AC005280, AC005548, AC005280, AC005651, AC005412, AC005881, AC004967, AF200465, AC006071, AC004590, AL136295, AC005519, AC004687, AL078634, AC005619, AC004687, AL096791, Z98752, Z8, AC008040, AC004815, AF064861, AC00564, AL049761, AC007993, AF038458, AP000692, AC005011, AC006468, AC006312, AL022165,	AL031	, AC003663, AC005670, AF11116
AC007066, AC005722, AL050404, AC AL133246, AC007298, AC005548, AC005280, AP000967, D86995, AC AL031575, AC002116, AC005412, AC007358, AC004967, AF200465, AC006071, AC004590, AL136295, AC006071, AC004590, AL136295, AC005519, AC004687, AL078634, AC005519, AC004687, AL078634, AC008040, AC004815, AF064861, AC008640, AC04148, AC004019, AC005664, AL049761, AC007993, AF038458, AP000128, AP000206, AC006261, AP006312, AL022165,	AL034	, AC005003, AC007371,
AL133246, AC007298, AC005548, AC005280, AP000967, D86995, AC, AL031575, AC002116, AC005412, AC007358, AC004967, AF200465, AC006071, AC003950, AL136295, AC007488, AC007546, AC010077, AC005519, AC004687, AL078634, AC008040, AC0046815, AF064861, AC008040, AC004148, AC004019, AC00564, AL049761, AC007993, AF038458, AP000128, AP000206, AC006468, AC006312, AL022165,	86582	AC007066, AC005722,
AC005280, AP000967, D86995, AC, AL031575, AC002116, AC005412, AC007358, AC004967, AF200465, AC005811, AC005811, AC005811, AC006071, AC004590, AL136295, AC00519, AC00519, AC004687, AL078634, AC008040, AC004815, AF064861, AC00564, AL049761, AC007993, AC006261, AP000692, AC005011, AC006468, AC006312, AL022165,	AC005	9, AL133246, AC007298, AC00554
AL031575, AC002116, AC005412, AC007358, AC004967, AF200465, AC00581, AC003950, AC006581, AC006071, AC004590, AL136295, AC007488, AC007546, AC010077, AC003071, AL096791, Z98752, Z8 AC003071, AL096791, Z98752, Z8 AC008040, AC004815, AF064861, AC005664, AL049761, AC007993, AF038458, AP000128, AP000206, AC006261, AP00692, AC005011, AC006468, AC006312, AL022165,	AP00C	, AC005280, AP000967,
AC007358, AC004967, AF200465, AC005881, AC003950, AC006581, AC006071, AC004590, AL136295, AC007488, AC007546, AC010077, AC005519, AC004687, AL0786481, AC003071, AL096791, Z98752, Z8, AC008040, AC004815, AF064861, AC007878, AC004148, AC004019, AC005664, AL049761, AC007993, AF038458, AP000128, AP000206, AC006261, AP00692, AC005011, AC006468, AC006312, AL022165,	AL034	, AL031575, AC002116, AC00541
AC005881, AC003950, AC006581, AC006071, AC004590, AL136295, AC007488, AC007546, AC010077, AC005519, AC004687, AL078634, AC003071, AL096791, Z98752, Z8, AC007878, AC004815, AF064861, AC005664, AL049761, AC007993, AF038458, AP00128, AP000206, AC006261, AP00692, AC005011, AC006468, AC006312, AL022165,	AC006	, AC007358, AC004967, AF20046
AC006071, AC004590, AL136295, AC007488, AC007546, AC010077, AC005519, AC004687, AL078634, AC003071, AL096791, Z98752, Z8 AC008040, AC004815, AF064861, AC007878, AC004148, AC004019, AC005664, AL049761, AC007993, AF038458, AP00128, AP000206, AC006261, AP006312, AL022165,	AL045	, AC005881, AC003950, AC00658
AC007488, AC007546, AC010077, AC005519, AC004687, AL078634, AC003071, AL096791, Z98752, Z8 AC008040, AC004815, AF064861, AC007878, AC004148, AC004019, AC005664, AL049761, AC007993, AF038458, AP000128, AP000206, AC006261, AP006312, AL022165,	AC007	, AC006071, AC004590,
AC005519, AC004687, AL078634, AC003071, AL096791, Z98752, Z8 AC008040, AC004815, AF064861, AC007878, AC004148, AC004019, AC005664, AL049761, AC007993, AC006261, AP000128, AP000206, AC006468, AC006312, AL022165,	AC006	, AC007488, AC007546,
AC003071, AL096791, Z98752, Z8 AC008040, AC004815, AF064861, AC007878, AC004148, AC004019, AC005664, AL049761, AC007993, AC006261, AP000128, AP000206, AC006468, AC006312, AL022165,	AL121	, AC005519, AC004687, AL07863
AC008040, AC004815, AF064861, AC007878, AC004148, AC004019, AC005664, AL049761, AC007993, AC006261, AP000128, AP000206, AC006468, AC006312, AL022165,	AC004	, AC003071, AL096791, Z98752, Z
, AC007878, AC004148, AC004019, , AC005664, AL049761, AC007993, , AF038458, AP000128, AP000206, , AC006261, AP000692, AC005011, , AC006468, AC006312, AL022165,	AC005	3, AC008040, AC004815, AF06486
, AC005664, AL049761, AC007993, , AF038458, AP000128, AP000206, , AC006261, AP000692, AC005011, , AC006468, AC006312, AL022165,	AC005	803, AC007878, AC004148, AC00401
, AF038458, AP000128, AP00020 , AC006261, AP000692, AC00501 , AC006468, AC006312, AL02216	ACOD5	, AC005664, AL049761, AC00799
4, AC006261, AP000692, AC00501 9, AC006468, AC006312, AL02216	AC002	, AF038458, AP000128, AP00020
9, AC006468, AC006312, AL02216	AC004	4, AC006261, AP000692, AC00501
	AC005	, AC006468, AC006312, AL022165
00212, AP000134, AF001549, Z99716,	APOOC	, AP000134, AF001549, Z99716, AL1
U82668, Z98051, AC002115, Z98884, AC00234	9	8, Z98051, AC002115, Z98884, AC00

5, Z83844, AC005207, AB 8, AL139054, AC004983,	ALO49764, ACO06057, AP000105, AP000037, AC006512, AC005480, AC000097, AC008115,	AC002072, AC005899, AC006942, Z73417, AF102137,	o`	, AC005358,	69, AC007216, AL021878,	AL079342, AL031729, AC006120, AP000354,	AP000210, AP000132, AC004797, Z94721, AL023284,	AB020873, AP000042, AP000110, AP000514,	AC007277, AL035460, AC005300, AL049712,	AC005932, AC005972, AP000350, AC007227,	5900, AC005953, AP000213	AC003037, AR036572, U91328, AC006013, Z85996,	, AC006344, AL121934	AL110502, AC005378, AC005544, AP000356	AW361997, AW362047, AW364154, AW362053,	, AI828567,	AW362017, AI559571, AI961395, AI922922,	8291, AI601204,	8715, AI922711, AI8	3816, AW150719, AI862126,	5885, AI870181, AI858824,	6673, AA555066, AW008365,	2031, AI923408, AI636160,	8713, AW150078, AI635039,	554505, AA582775, AI564401, AW177640,	151832, AI433333, AI274367, AA418161,	955997, AI640719, AI625261, AW191047,	357830, AI588841, AI721066, AI858311,	AI683781, AW361718, AI581904, AI373075,	AI628535, AW364741, AI564624, AI587610,	AA652197, AA553882, AI859018, AW193154,	AI858269, W52120, AI445544, AW062541, AI659775,	859238, AI417458, AW338495, AW298327,
AD	AL	AC	AC	AC	AC	AL	AP	AB .	AC	AC	AC003	AC	AC	AL	Preferably excluded from the AW	on are one or more	leotides comprising a	ence described by	a-b, where a	eger between 1 to 2863 of	integer of	υ.	)f	residues shown in SEQ ID	NO:122, and where b is greater than AI5	or equal to a + 14.	AI95	AI85	AI	AI	AP	AI	AI
						_,									722258																		
															HUFAC36																		
															122																		

AW364615, AW364573, AW351747, AI799729,
, AI96377
, AW375899, AI799381, AI27597
141, AW351538, AW364165,
867651, AI648422, AI422584, AA37695
AA366895, AA377282, AI669320, AI811793,
 548, AA295047, AA417910,
, AI833145, AI824496, T34
AI982524, AI8
, AI648451, AW376236, AA53
76235, AI933787, AW366558,
01121, AW364616, AI659355, AW364
37, AI866055, T24880,
0, T24837, T35699, AI418084,
), AW375551,
2, AI540524, AI568068, C06052,
), T24449, AI553666,
AI721090, AI973152, T25155, Al
), AW364785, AI860027,
8, AI690813, AI540674, AI538
17, AI918677, AI796743, AW10543
1, AW194014, AL045496,
27233, AA937566, AI873550,
5, AI537191, AI559752,
18809, AI699865, AI445829, AI342710
081740, R40363, AW083573, AI500658, AI63
590043, AI440238, AW084396,
 9, AI621341, AW029216,
AI863002, AW105460, AW084896, AI539260,
l, I68732, AF026816, AF090903, D8303
X8
AL023657,
AF151685, AF017437, I89947, AL137537, AL049938,
AF185614, AR038854, AL050277, X06146, AF124396,
 , AL133665, A60094, AF0315
A77035, U87620, E12747, A21103, E03671,

				ARU68466, ALIL/438, AF040/23, U38419, AF060555,
				3338, AFUSUNS, AFZUN464, 3633 - AFUSUNSON 1186379 - T2
				281, AF161413, AL137550, AF167995, A08
				, X93328, AF008439, A08911, A65341,
				E12580, A93914, AL133047, I32738, X68497,
				AJ001388, A15345, A26498, E128
				,
				A58545, AR050959, AL133067,
				0340,
				309, AL049276, AL122103
				AF109683, I33392, AL117460, Z97214, A18777,
				AF
				X72624, AF111849, U31501, AR034821, AF032666,
				A12522, AF111112, AF098484, AR066485, A76335,
				AL137554, AL110221,
				AB031064, AL080146, U57715, Z82022, AL133075,
				AF124728, AL137284, AF029750, AL133031,
				AF061943, X82397, AL109672, AL117626, AL049423,
	_			55
123	HHFHB49	723136	Preferably excluded from the	AA054421, T24430, AW206410, AI660910, AC004202,
			present invention are one or more	AP000518, AB023054, Y07828
	_		tides comprising a	
	_		ednence des	
	_		the general formula of a-b, where a	
	_		eger between 1 to 667	
	_		SEQ ID NO:123, b is an integer of	
-	_		15 to 681, where both a and b	
	_		to the positions of	-
			nucleotide residues shown in SEQ ID	
•			NO:123, and where b is greater than	
			or equal to a + 14.	
124	HFIBH05	725110	Preferably excluded from the	AA625451, AI089287, AA282874, AA398984, H72493,
			present invention are one or more	AA137263, AI434776, N33821, AA482849, H79114,

			ides comp	R23405, R25093, AI983837	
			nucleotide sequence described by the general formula of a-b, where a		
			()		
-		- · · -	SEQ ID NO:124, b is an integer of		
			15 to 606, where both a and b		
			correspond to the positions of		
	_		nucleotide residues shown in SEQ ID		
			ਲ		
			or equal to a + 14.		
125	HKIAA57	725201	Preferably excluded from the	AI741792,	, AI620910,
			present invention are one or more	AW027395, AI799988, AA659728,	, AA883923,
			polynucleotides comprising a	AI361118, AI694798, R39993,	R39993, AI421599, AI421231,
			nucleotide sequence described by		AA166749, R73993,
_			the general formula of a-b, where a	AA306989, D59334, W21931, AA	W21931, AA918493, R73900,
			is any integer between 1 to 1197 of	AA059363, AI368574, T10593,	AA476990, R43798,
		.,	SEQ ID NO:125, b is an integer of	AA410954, T10567, AB020676	
		. ,	15 to 1211, where both a and b		
			correspond to the positions of		
			shc		
			NO:125, and where b is greater than		
			or equal to a + 14.		
126	HRKAB52	726122	Preferably excluded from the	, AI733029,	
			present invention are one or more	1, AI288746,	, AI318112,
			polynucleotides comprising a	AI301846, AI304527, AI274206,	, AA878571,
			nucleotide sequence described by	_	, AW075794,
			the general formula of a-b, where a		T68446, AA732362, T93796,
			is any integer between 1 to 867 of	AI720888, AA906537, AI681040,	, Z79996, AC002394,
			SEQ ID NO:126, b is an integer of	AC006059, Z95126, AC007198, AL022336,	AL022336, AC007065,
			15 to 881, where both a and b	AC004220, AC002980, AL024493, AC005549	, AC005549,
			correspond to the positions of	U91325, AL021182,	AC007380, AC005228,
			nucleotide residues shown in SEQ ID	Z82205, Z83826, AC004551, AC	003119, AF039905,
			NO:126, and where b is greater than	AP000292, AP000043, AP000111, AL008626	, AL008626,
			or equal to a + 14.	AC002992, U80460, AC004088	
127	HPCAN95	727365	Preferably excluded from the	AA007664, AI803958, AI167454,	, AI968968,
	-		present invention are one or more	AI247561, AA148881, AA007627	, AI375078, W23292

			nolymnaleotides comprising a	
			al formula	
			or a z, micro	
			1000t	
			:12/, D 15 all lilleger	
			where both a and	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:127, and where b is greater than	
			or equal to a + 14.	
128	HCQCV54	729143	Preferably excluded from the	N45700, H63509, H54749, AA789241, AI073405,
			present invention are one or more	AL137699
			polynucleotides comprising a	
			nucleotide sequence described by	
•			the general formula of a-b, where a	
			SEQ ID NO:128, b is an integer of	
			15 to 1287, where both a and b	
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:128, and where b is greater than	
			l to a + 14.	
129	HLJEA54	729231	Preferably excluded from the	AI079148, AA532656, AP000548, AL031120, AP000365
			present invention are one or more	
			polynucleotides comprising a	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			is any integer between 1 to 589 of	
			SEQ ID NO:129, b is an integer of	
			15 to 603, where both a and b	-
			correspond to the positions of	
			nucleotide residues shown in SEQ ID	
			NO:129, and where b is greater than	
			or equal to a + 14.	
130	HTWCR70	731881	Preferably excluded from the	AI246796, AA847499, AL047080, N22516, AA504694,
			present invention are one or more	T40848, R91796, AA492015, AA503468, N70293,

	polynucleotides comprising a	AW024018,	AW028943,	AI886629,	F04766, AL038901,
	sedne	AI249688,	AI491828,	AL038533,	N23846, AI963045,
	$\neg$	AA515138,	AW274182,	AI916406,	AW302711,
	eger between	AA535937,	AW439480,	AA488746,	AA515727,
	SEQ ID NO:130, b is an integer of	AJ246003,	AC006241,	AC007421,	AL121658, U91323,
	ø	AL109623,	AC002558,	AL096701,	AL009181,
	correspond to the positions of	AC002551,	AC006449,	AC005480,	AC007055,
		AC004382,	AL035086,	AC007283,	AC004263,
	NO:130, and where b is greater than	AC007546,	AC002091,	AC000379,	AC003101, U95742,
	or equal to a + 14.	AC005527,	AL139054,	AC005399,	AL049829,
		AC004983,	AC005529,	AL050307,	AP000553,
		AF196969,	AC005488,	AC005225,	AL133245,
		AL049576,	AC005015,	AC007225,	AL022165,
		AC004638,	AC004703,	AL031432,	
		AC000353,	AC000025,	AC007050,	U62293, AC007637,
		AC002565,	AL080243,	AL121603,	AC005231,
		AL049830,	AF109907,	AC006537,	AP000692,
		AC005288,	AC004131,	AC005180,	Z95113, AL022476,
		AF196779,	AC002312,	AL109758,	AC005920,
•		AC016025,	AC005914,	AL031588,	AC004408,
		AC006120,	AC004686,	AL096791,	AC004491,
		AL009183,	AC004858,	AF134726,	AC004383,
		AC005520,	AC004859,	AC004813,	AL049843, Z98941,
		AP000212,	₹#	σ	Z83840, AC006211,
		AC006960,	U96629, A	AC005091, A	AF001549, AC002300,
		AC006157,	AC002288,	AC004883,	Z98884, AC007151,
		AC004953,	AL031680,	AC005933,	
		AC007666,	AL079342,	AC002470,	
		AL035413,	AC005257,	AP000030,	
		AC007227,	AL031985,	AC004257,	AP000248,
		AC006088,	AL031577,	Z82244, AL022318,	L022318, AC004999,
		AC002996,	AC005823,	AC005972,	AC005280, U80017,
		AC002070,	AC004820,	AL023807,	AC006530,
		AC007226,	AC002310,	AL049692,	
		m	AL133163,	AL031283,	AC007114,
		AL049776,	AC006111,	AL109613,	AC004019,

				AT.031984 AC006312 AF053356 AC007371 Z98051
				548, Z97055, AC006141, AP000141, AC00570
				, AL078638, AC005632, Z95152,
				, AC005839, Z82201, AC005740,
_				AF001552, AL049872, U82668, AC005207, AC005412,
				AP000065, AC002045, AC004812, AC004134,
				AC006121, AC005264, AF165926, AC002477,
				AC004106, AC004531, AL050341, AL022238,
_				AC004583, AC006538, AF111168, AC004217,
				AC005037, AC005829, AC007124, Z84469, AF067844,
				L035420, AL
				AC004955, AC005082, AC006501, AC000159,
	_			AC004596, AC005484, AC005210, AC006441,
				AC002059, AL031311, AC004098, AL135744,
				AC006254, AC010205, AL035461, AC000004,
	-			AC004991, AF139813, AC012384, AC007370,
				AC005531, AL049780, AC005874, AF134471,
				AC016830, AL031685, AL021546, AL035400,
				AC005776, AC003002, AC002400, AL132642,
				ın
				A
				AL049869,
				AC000052, AL022323, AC003071, AP000211,
				26,
				AC009721, AP000555
131	HSXDD55	732280	Preferably excluded from the	H19388, AA121710, H12126, AA429913, AA446069,
			present invention are one or more	AW104301, AB002349
	•		tides comp	
			nucleotide sequence described by	
			the general formula of a-b, where a	
			eger betwe	
	-		SEQ ID NO:131, b is an integer of	
			15 to 776, where both a and b	
			correspond to the positions of	-
			ide	
			NO:131, and where b is greater than	

			Or Agial to a + 14	
135	HAGEX59	735603	preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 1394 of SEQ ID NO:135, b is an integer of 15 to 1408, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:135, and where b is greater than or equal to a + 14.	AI138441, AI400746, AI400746, AIA60767
136	HAVMG19	739061	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 888 of SEQ ID NO:136, b is an integer of 15 to 902, where both a and b correspond to the positions of nucleotide residues shown in SEQ ID NO:136, and where b is greater than or equal to a + 14.	AA043107, AA460757, AA808519, AA482341, AW088728, AL041651, AI305256, AA885398, AI808361, D53182, AI689925, AI123220, N99552, AN130266, AA861771, AA040860, AI278439, AA134816, W87524, W89049, AI301074, AA927150, W87525, AI086181, AI683247, AI633628, AI301808, AW236826, AA339816, AA649134, AA806264, AI078052, AA460158, AA649134, AA806264, AI078052, AA460158, AA909561, R46365, R54321, AI472152, T23855, AI769853, AW074642, W57681, AA039402, AW072224, T80969, AA991845, AI472163, AI831540, AI686045, AA716427, AA029761, T97173, AI738802, T81108, R67010, AA482244, W89104, R49077, T40888, AI583709, N71544, AB028951, AL122055
137	HLEAL50	741134	Preferably excluded from the present invention are one or more polynucleotides comprising a nucleotide sequence described by the general formula of a-b, where a is any integer between 1 to 716 of SEQ ID NO:137, b is an integer of 15 to 730, where both a and b correspond to the positions of	AA745961, AII10640, AA581433, AW419403, AL119355, AA843874, AA309370, AI439860, AI887321, AW130042, AA174085, AII10720, AL133942, AW177226, AA601264, AW177120, AI818151, AW150375, AA152091, AW089864, AI925647, AW157413, AI983921, AI927861, AI367384, AI858607, AA493735, AW089655, W03759, H97952, AI811494, AW190486, AI761656, AI685116, AW167319, AI610776, AA679519, AI688902,

	nucleotide residues shown in SEQ ID	AI732743, AA493998, AL137994, AI872415,
	NO:137, and where b is greater than	AI264673, AI334099, AAI36637, AA659014,
	0 a + 14.	9, AA189081, AA130536, AA77335
	•	AI924175, AA767353, AA631430, AI874256,
		AW177231, AW235478, N24958, AI088796, W49501,
		7, AW151
		AI749571, AI376984, AI817158, AA778304,
		ω,
		AI819528, AI568919, AI862874, AL041411,
		AW074001,
		AA932087,
		, AI479035, AA427754
		AA121916, AA130476,
		AI733728, AA663566, W87732
-		N792
		AL041417, R80440, AI250812
		3, AI346802, AW242735, AI5911
		AA868708,
		53572, AI499286, AW004844,
		, AI114529, AA346162, AA17635
		33889, AA501873, AA807609,
		877, AA709024, AI557354,
		, AA911409, AA135303
		, AI246569, H12832, AA946637
		AI453790, H91008, AI110627, AA65
		43039, AA081993, AI025602, AI272961, T4
		281, H73189, AA854527, AI56
		1, AI632138, AA889273, AI5
		198, AW102963, AI095849, AA363058
		06932, AA
		2, AA771730
		AC008394, AL079352, AL031663, AL136018,
-		9, U20230, AL049794,
		K00628, AC003082, AC004467, AL049561, AC005194,
		L29074, AC009320, AP000687, AJ229041, AC005138,

1 1117339 AC005034 AL031903 AC004070
4, Z80107, AL022401, AC
951, AC003081, AC007671
939, AC005016,
AC006070, Z75741, AC
AC007372, AC005509, AC008170, Z81001, AC010168,
AC006561, Z76735, AC004750, AL
Z83827, AC010382, AC004647, AF020802, AL009173,
, AC004957, AC006484,
9, AC006578,
AC004454, AL121578, AC006371, AF064864,
AC006197, AL033403, AC006037, AF064860,
, AC00405
AL034369, AC000377, AL133512, AL049837,
AC005045, AC006971, AC000114, AL009172, Z99495,
Z82899, AC
, AL133381, AC004538
, Z96810, AL009174,
6, AC003016, Z93928, AC00495
, Z83819, AL033521, AL
', AL049546,
, AC006946, AP000473, AL031655
, AL021068, AC006144,
1, AC002524, AL049564, AB033055,
5, AC010849, AC007110
AL049828,
AC005352, AC004911, AL078474, Y10196, AL121782,
AC006316, AL109662, AL133249, AL009176,
AL049691, AF188025, AC004535, AC006455,
AC007090, AC002367, AJ006343, AC004820,
AC004907, AL008710, AL034408, AL031673,
_
AL022151, AC005221, AC007450, AC005017,
AC005066, AP000474, AC004869, AC004415,

				0711000	0501C11V	21130004
				ACU # 100'	7, ALLEIO30,	, 211000
				AC006324,	AL022577, AC007402, AC	AC004674,
				AJ272265,	AC002526, AL022146, AC	AC002452,
				AC006504,	AC004082, AC005184, AC	AC008009,
				AL033379,	Z72001, AF002991, AC00	AC002478, AF165175,
				AC005201,	, Z70232,	AC002070, AC004949,
				AC004831,	AL049642, Z94055, Z700	Z70049, AP000014,
				AC004128,	AC007971	AL117326, U80460,
				AC000111,	AL049710, AC003693, AL	AL096773,
				AP000454,	AC005873,	AP000127,
				AP000205,		AP000948, Z84720,
				AL034561,		AC006002, AC007535,
				AL035427,	Z92545, AL049176, AL03	AL031782, AL031586,
				AC005271,	5,	AL079303,
				AP000244,	, AC007628,	AC004130,
				AL109753,	Z69649, AP000496, AB02	AB023054, AC004677
138	HCPAC07	741257	Preferably excluded from the	AA729528,	AI659354, H92153, AI60	AI609659, AC007842
			present invention are one or more			
			polynucleotides comprising a			
			nucleotide sequence described by			
			the general formula of a-b, where a			
			SEQ ID NO:138, b is an integer of			
			15 to 524, where both a and b			
			d to the po			
			ide residues sho			
			NO:138, and where b is greater than			
			or equal to a + 14.			
139	HOSEQ61	741804	Preferably excluded from the	AI191412,	3, AW188995,	AI817093,
			present invention are one or more	AI682959,	AI684692,	AI921724,
			polynucleotides comprising a	AI742357,	AI920893,	AI056062,
			nucleotide sequence described by	AW080580,	٠,	AI399967,
			the general formula of a-b, where a	AI288672,	N47530, AI025387, R374	R37474, AI399768,
			is any integer between 1 to 855 of	AI082088,	_	AA992114,
			SEQ ID NO:139, b is an integer of	AW172410,	AA804760, AI277609, AI	AI419244,
			15 to 869, where both a and b	AI581273,	AW264450, AI690471, AI	AI201792,